

SurgWeek-2 Appendices

Appendix A: List of excluded procedures

The SurgWeek-2 study will include all operations (elective or emergency) performed in an operating theatre, excluding minor procedures. All surgical specialties are included. Both day case surgery and inpatient surgery are included. Both children and adults are included. The minor procedures that are excluded are listed below.

Specialty	Excluded procedures
Abdominal surgery	Ascitic drain (drainage of peritoneal cavity)
	Endoscopic ultrasound
	Laparoscopic ultrasound
Breast surgery	Breast biopsy
Cardiac surgery	Insertion of cardiac pacemaker
	PCI: percutaneous coronary intervention
	Transluminal balloon angioplasty of coronary artery
Colorectal surgery	Colonoscopy (diagnostic or therapeutic)
	Flexible sigmoidoscopy (diagnostic or therapeutic)
	Proctoscopy (diagnostic or therapeutic)
Dental procedures	Implantation of tooth
	Insertion of dental prosthesis
	Orthodontic operations
	Restoration of tooth
	Extraction of tooth
Gynaecology	Cervical biopsy
	Colposcopy (diagnostic or therapeutic)
Obstetrics	Any vaginal delivery (normal delivery, breech delivery, forceps delivery, vacuum delivery)
	Repair of obstetric laceration

Ophthalmology	Removal of foreign body from cornea
Orthopaedics	Bone biopsy
	Injection into joint
	Muscle biopsy
Otolaryngology	Nasendoscopy (diagnostic or therapeutic)
	Packing of cavity of nose
Thoracic surgery	Bronchoscopy (diagnostic or therapeutic)
	Chest drain
Upper gastrointestinal surgery	ERCP: endoscopic retrograde cholangiopancreatography (diagnostic or therapeutic)
	Liver biopsy
	OGD: Oesophago-gastro-duodenoscopy (diagnostic or therapeutic)
Urology*	Bladder biopsy
	Extracorporeal shock wave lithotripsy (ESWL)
	Flexible cystoscopy (diagnostic or therapeutic)
	Percutaneous nephrostomy
	Percutaneous nephrolithotomy (PCNL)
Vascular surgery	Transluminal (endovascular) procedures on arteries (diagnostic or therapeutic), including with open cut down to the artery
	Transluminal (endovascular) procedures on veins (diagnostic or therapeutic)
Other	Insertion of central venous catheter/ line (CVC)
	Lumbar (spinal) puncture
	Percutaneous tracheostomy
	Skin biopsy (including shave biopsy of skin)
	Therapeutic epidural injection

*Note: transurethral resection of the prostate (TURP), transurethral resection of bladder tumour (TURBT), rigid cystoscopy under general anaesthesia, and insertion of ureteric stent should be included.

Appendix B: Case Report Forms (CRF)

In the CRF the sections in grey will become available via branching logic if triggered by prior data entry.

Demographic and pre-operative data		
1.1	Period operated in	1, Period 1 (00:00 31 st August 2026 – 23:59 6 th September 2026) 2, Period 2 (00:00 7 th September 2026 – 23:59 13 th September 2026) 3, Period 3 (00:00 14 th September 2026 – 23:59 20 th September 2026) 4, Period 4 (00:00 21 st September 2026 – 23:59 27 th September 2026) 5, Period 5 (00:00 28 th September 2026 – 23:59 4 th October 2026) 6, Period 6 (00:00 5 th October 2026 – 23:59 11 th October 2026)
2.1	Sex	1, Female 2, Male
2.2	If female and age 10-59: pregnancy status	1, No 2, Yes – first trimester (<13 weeks) 3, Yes – second trimester (13-28 weeks) 4, Yes – third trimester (>28 weeks)
3.1	Age	0, 0-4 weeks 1, 5 - 52 weeks 2, 1-4 years 3, 5-9 years 4, 10-17 years 5, 18-29 years 6, 30-39 years 7, 40-49 years 8, 50-59 years 9, 60-69 years 10, 70-79 years 11, 80-89 years 12, 90 years or more
4.1	ASA	1, Grade I 2, Grade II 3, Grade III 4, Grade IV 5, Grade V 6, Grade VI
5.1	If aged 18 years or more: BMI	Number
	Link to calculator: https://www.bupa.co.uk/health-information/bmi-calculator	
5.2	If aged less than 18 years: Weight (kg)	Number
6.1	Co-morbidities <i>Please tick all that apply</i>	1, Ischaemic heart disease 2, Congestive heart failure 3, Cerebrovascular disease 4, Diabetes mellitus – insulin dependent 5, Diabetes mellitus – non-insulin dependent 6, Chronic kidney disease (baseline creatinine >176.8 µmol/L or >2 mg/dL) 7, Human immunodeficiency virus (HIV) 8, Hypertension 99, None of the above
6.2	If HIV: HIV status	1, Untreated HIV 2, HIV with CD4 cell count <200 cells/mm ³ 3, HIV with CD4 cell count 200-500 cells/mm ³ 4, HIV with CD4 cell count >500 cells/mm ³ 5, HIV with CD4 cell count unknown
7.1	Clinical frailty score Link: https://www.bgs.org.uk/sites/default/files/content/attachment/2018-07-05/rockwood_cfs.pdf	1, 1 – Very fit 2, 2 – Well 3, 3 – Managing well 4, 4 – Vulnerable 5, 5 – Mildly frail 6, 6 – Moderately frail 7, 7 – Severely frail – completely dependent for personal care 8, 8 – Very severely frail 9, 9 – Terminally ill
8.1	Smoking status <i>Please tick all that apply</i>	1, No – never smoked 2, No – ex smoker, stopped ≥ 6 weeks ago 3, No – ex smoker, stopped in the last 6 weeks 4, Yes – current smoker 5, No, never vaped 6, No – previously vaped, stopped ≥ 6 weeks ago 7, No – previously vaped, stopped in the last 6 weeks 8, Yes – currently vapes 9, Unknown smoking or vaping status
9.1	Type of admission	1, Elective (surgery on a planned admission) 2, Emergency (surgery on an unplanned admission)
10.1	Indication for surgery (broad category)	1, Benign 2, Malignant / pre-malignant 3, Trauma 4, Obstetrics
10.2	If malignant: Intent of this operation	1, Curative 2, Palliative
10.3	If malignant: Extent of malignancy	0, Pre-malignant 1, Primary only 2, Nodal metastases 3, Distant metastases 4, Not staged
10.4	If trauma: Mechanism of injury <i>Please tick all that apply</i>	1, Fall less than 2 metres 2, Fall more than 2 metres 3, Road traffic collision – occupant 4, Road traffic collision – pedestrian 5, Road traffic collision – cyclist 6, Gunshot 7, Stab 8, Other penetrating injury (not gunshot/stab) 9, Assault – blunt force 10, Blast / explosion 11, Crush injury 12, Industrial / Occupational 13, Animal bite 14, Sport related 15, Burns or thermal injury 16, Self harm 17, Other, please specify
10.5	If trauma: Identify all of the injuries <i>Please tick all that apply</i>	1, Single acute fracture 2, Multiple acute fractures 3, Hand injury 4, Traumatic brain injury 5, Skull fracture 6, Facial fracture 7, Neck injury (vascular/airway) 8, Rib fractures 9, Pneumothorax/Haemothorax 10, Lung contusion 11, Cardiac injury 12, Abdominal solid organ injury 13, Abdominal hollow viscus injury 14, Vertebral fracture 15, Spinal cord injury 16, Upper limb dislocation 17, Upper limb soft tissue injury 18, Lower limb dislocation 19, Lower limb soft tissue injury 20, Vascular injury 21, Nerve injury
11.1	Body region of main operation	1, Head and neck 2, Thorax 3, Abdomen and Pelvis 4, Upper limb 5, Lower limb 6, Spine
12.1	Indication for operation (specific diagnosis)	See list
9.2	If emergency admission: On presentation to hospital, was the patient septic?	0, No 1, Yes
9.2a	If sepsis: Duration of time from identification of suspected sepsis in hospital to first antibiotics being given	1, <30 minutes 2, 30-59 minutes 3, 60-119 minutes 4, 120 minutes - 24 hours 5, No antibiotics given within 24 hours
9.3	If emergency admission: On presentation to hospital, were there signs of shock?	0, No 1, Yes
9.4	If emergency admission: Time of symptom onset or injury to presentation to hospital	1, Under 12 hours 2, 12-23 hours 3, 24-47 hours 4, 48-71 hours 5, 72 hours or more
9.5	If emergency admission: Time from presentation to hospital to review by surgical team	1, <1 hour 2, 1-2 hours 3, 3-6 hours 4, 7-12 hours 5, 13-24 hours 6, Over 24 hours
9.6	If emergency admission: Time from review by surgical team to decision for operation	1, <1 hour 2, 1-2 hours 3, 3-6 hours 4, 7-12 hours 5, 13-24 hours 6, Over 24 hours
9.7	If emergency admission: Duration of time from decision to operate (booking of theatre case) to patient arrival in theatre	1, <2 hours 2, 2 - 6 hours 3, 7-18 hours 4, >18 hours
9.8	If emergency admission: What was the intended window from decision to operate (booking the theatre case) to the operation?	1, <2 hours 2, 2 - 6 hours 3, 7-18 hours 4, >18 hours 5, No window set 6, Unknown
9.9	If emergency admission: Immediately prior to attending this hospital, did patient seek care from anywhere else? <i>Please tick all that apply</i>	0, No 1, Yes – Traditional or faith healer 2, Yes - Community health worker 3, Yes – Family or primary care doctor 4, Yes - Another hospital transferred this patient to this hospital 5, Yes – Other, please specify 6, Unknown

9.9a	If transfer from another hospital: What was the reason for hospital transfer? <i>Please tick all that apply</i>	1, For input from a surgical speciality not available at referring hospital 2, No surgeon at referring hospital 3, Anaesthetic requirement 4, No operating theatre 5, Critical care (High dependency or intensive care unit) requirements 6, Diagnostics; for example no CT scanner 7, Interventional radiology unavailable 8, Blood product availability 9, Equipment limitations (robot/laparoscopy/stapling devices etc) 10, Complexity requires tertiary centre (i.e. patient factors) 11, External disruption; for example power issues, flooding, staff strikes 12, Financial constraints 13, Other, please specify
9.10	If elective admission: How many days passed between the decision to operation (or patient being booked for the operation) to having the operation?	Integer (days)

Intra-operative		
13.1	Was the WHO Safer surgery checklist used in theatre?	0, No 1, Yes
14.1	Grade of the most senior surgeon in theatre	1, Consultant or equivalent 2, Doctor/surgeon with 3 or more years of experience 3, Doctor/surgeon with 1-2 years of experience 4, Non-physician
15.1	Grade of primary surgeon (Defined as performing over 50% of the critical operative steps)	1, Consultant or equivalent 2, Doctor/surgeon with 3 or more years of experience 3, Doctor/surgeon with 1-2 years of experience 4, Non-physician
16.1	Grade of the most senior anaesthetist in theatre	1, Consultant or equivalent 2, Doctor with 3 or more years of experience 3, Doctor with 1-2 years of experience 4, Non-physician
17.1	Type(s) of anaesthesia <i>Please tick all that apply</i>	1, General (gaseous) 2, General (total intravenous) 3, Neuraxial - epidural 4, Neuraxial - spinal 5, Regional nerve block 6, Sedation 7, Local anaesthetic 8, None
18.1	Time of knife to skin (start of operation)	1, 08:00-17:59 2, 18:00-23:59 3, 00:00-07:59
19.1	Main operation	See drop down list
20.1	Planned initial operative approach	1, Open 2, Standard minimally invasive (e.g. laparoscopic, thoracoscopic or equivalent video-assisted technique excluding robotic assisted) 3, Robotic assisted 4, Hybrid (minimally invasive and open) 5, Endoluminal 6, Natural orifice transluminal
21.1	Final (completion) surgical approach	1, Open 2, Standard minimally invasive (e.g. laparoscopic/thoracoscopic or equivalent video-assisted technique excluding robotic assisted) 3, Robotic assisted 4, Hybrid (minimally invasive and open) 5, Endoluminal 6, Natural orifice transluminal
22.1	Degree of intra-operative contamination	1, Clean 2, Clean-contaminated 3, Contaminated 4, Dirty
23.1	Did the patient receive antibiotics at induction of anaesthesia?	1, Yes, at induction of anaesthesia 2, No, but had a dose within the previous 8 hours 3, No and did not have any antibiotics in the preceding 8 hours.
24.1	How was the main surgical wound closed?	1, Primary closure 2, Left to heal by secondary intention 3, No external surgical wound or not applicable
24.2	If surgical wound (i.e. 24.1 is 1 or 2): What dressing was used for the main surgical wound?	1, Antimicrobial or bacteria-binding (e.g. iodine impregnated, silver impregnated, chlorhexidine impregnated or hydrophobic fibre/DACC) 2, Closed incision negative pressure (ciNPWT) (canister-based or single use) 3, Glue-containing (e.g. cyanoacrylate skin adhesive, alone or with self-adhering mesh) 4, Simple dressing only (e.g. gauze, adhesive island dressing, transparent film) 5, No dressing
24.3	If wound healing by secondary intention: Location of the wound left to heal by secondary intention	1, Foot 2, Leg 3, Abdomen 4, Other, please specify
25.1	Skin preparation <i>Please tick all that apply</i>	1, Chlorhexidine in alcohol 2, Aqueous chlorhexidine 3, Povidone-iodine (aqueous) 4, Povidone-iodine in alcohol 5, Alcohol only 6, Olanexidine 7, Ioban drape or equivalent 8, Other, please specify 9, Unknown
26.1	Were there any power outages during the operation?	0, No 1, Yes, 1-14 minutes total power outage 2, Yes 15-29 minutes total power outage 3, Yes, 30-59 minutes total power outage 4, Yes, > 1-hour total power outage 5, Unknown
26.2	If there was a power outage: Did this result in any change of procedure?	0, No 1, Conversion from minimally invasive to open 2, Abandoned procedure objective 3, Alternative procedure performed 4, Other, please specify
26.3	If there was a power outage: What was the impact to the patient?	1, No impact 2, Minor complication 3, Moderate complication 4, Life-threatening complication

Post-operative data		
27.1	Has this patient had follow-up within 30 days? <i>Please tick all that apply</i>	0, No follow-up in 30 days, data collected from hospital record review 1, Yes - In-person clinic follow-up 2, Yes - Telephone or video call
27.2	If follow-up in person or telephone/video: On what post-operative day did the patient return to normal activity? (This could include work, care or community roles) (If the patient has not returned to normal activity, please enter 99)	Integer (days)
28.1	30-day Clavien-Dindo grade	1, Grade I 2, Grade II 3, Grade III a 4, Grade III b 5, Grade IV a 6, Grade IV b 7, Grade V 99, No complications
28.2	If 30-day complication: Which post-operative complication(s) did the patient have? <i>Please tick all that apply</i>	1, Surgical site infection or fracture-related infection 2, Deep organ space infection 3, Pneumonia 4, Unexpected ventilation 5, Acute respiratory distress syndrome (ARDS) 6, Pulmonary embolism 7, Deep vein thrombosis 8, Pulmonary aspiration or aspiration pneumonitis 9, Urinary tract infection 10, Central line infection 11, Sepsis or septic shock 12, Myocardial infarction 13, New acute heart failure or clinically significant cardiac arrhythmia requiring treatment 14, Stroke 15, Postoperative haemorrhage requiring transfusion 16, Anastomotic leak 17, Acute kidney injury resulting in dialysis 18, Wound dehiscence 99, None of the above
29.1	30-day reoperation	0, No 1, Yes, planned re-operation 2, Yes, unplanned re-operation 3, Yes, both planned and unplanned re-operations
30.1	Post-operative length of stay (days) (Day 0 is the day of operation)	Integer (days)
31.1	Was the patient admitted to critical care postoperatively (high dependency unit or intensive care unit)? <i>Please tick all that apply</i>	0, No - not required 1, No - planned to go to critical care but no bed available 2, Yes - planned from theatre 3, Yes - unplanned from theatre 4, Yes - unplanned from ward
31.2	If critical care admission: On how many postoperative days, within 30 days of the operation, did the patient spend in a critical care unit? (day 0 is the day of the operation)	Integer (days)
32.1	Did your patient have an enhanced recovery after surgery (ERAS) protocol or equivalent?	0, No 1, Yes
33.1	If malignancy (Q10.1 = malignant/pre-malignant): Was histology performed?	1, Not performed 2, No specimen 3, Yes – benign 4, Yes – malignant with R0 resection margin 5, Yes – malignant with R1 resection margin 6, Yes – malignant with R2 resection margin 7, Yes – malignant but no resection margin provided
34.1	On how many postoperative days, within 30 days of the operation, did the patient receive at least one dose of antibiotics	Integer (days)
35.1	Did the patient have a wound swab sent for microbiology?	0, No 1, Yes, but no growth on any specimens 2, Yes, growth on one or more specimens but organisms with no antimicrobial resistances 3, Yes, growth of organisms in one or more specimens with antimicrobial resistance 4, Yes, growth but no sensitivity performed or available
35.2	If 35.1 is option 3: which organisms had resistance to antibiotics? <i>Please tick all that apply</i>	Enterobacteriales: 1, Escherichia (e.g. E. coli) 2, Klebsiella (e.g. K. pneumoniae) 3, Enterobacter 4, Proteus Specific resistances: 5, ESBL-producing Enterobacteriales (e.g. E. coli, Klebsiella) 6, Carbapenem-resistant Enterobacteriales (CRE) 7, Pseudomonas aeruginosa (multidrug-resistant) 8, Methicillin-resistant Staph aureus (MRSA) 9, Vancomycin-resistant Enterococcus (VRE) Anaerobes: 10, Bacteroides fragilis 11, Other Bacteroides spp. 12, Peptostreptococcus 13, Clostridium spp. Other: 14, Gram negative 15, Enterococcus 16, Other not listed, please specify
36.1	30-day hospital re-admission	0, No 1, Yes
36.2	If re-admitted: Length of readmission stay (If discharged on the same day as readmission then enter 0. If multiple readmissions within 30 days, please state the total number of readmitted days)	Integer (days)
37.1	How was the majority of the cost of surgery supported?	1, Insurance provided by the government (national or regional level) 2, Insurance provided by employer (or household members' employer) 3, Insurance that the patient has privately arranged and paid for 4, Insurance but unknown how this was arranged 5, External funds or grants awarded by charities/NGOs 6, Out-of-pocket payments (patient paid the hospital directly) 7, Free at the point of use (e.g. UK National Health Service)
37.2	If option 1-5 from 'how was the majority of the cost of surgery supported': Did the patient pay anything out of pocket?	0, No 1, Yes 2, Unknown
37.3	If 37.2 is yes or 37.1 is 6: Did an inability to pay result in a delay to the patient having an operation?	0, No 1, Yes 2, Unknown

If operative approach is robotic (branch from question 20.1 and Question 21.1)	
Robotic platform	1, da Vinci Xi 2, da Vinci X 3, da Vinci Si 4, da Vinci 5 5, Versius 6, Hugo RAS 7, Senhance 8, Moon 9, KangDuo or KD-Surgibot 10, Tourmal/Rebo 11, Hinotori 12, ROSA robotic system 13, Sina surgical system 14, Microport 15, Sentire surgical system 16, Mako robotic-arm assisted surgery system 17, Ottava surgical system 18, SSi Mantra 19, Single-port system, please specify 20, Other, please specify
If no robot had been available today, what would have been done for this patient?	1, Cancel this operation 2, Open surgery 3, Laparoscopic or other minimally invasive equivalent surgery
Was team briefing completed, including plans for emergency undocking and system failure?	0, No 1, Yes
Lifetime number of robotic cases for the console surgeon	1, 0 2, 1-10 3, 11-25 4, 26-50 5, 51-100 6, Over 100
Grade of the bedside assistant	1, Consultant or equivalent 2, Doctor/surgeon with 3 or more years of experience 3, Doctor/surgeon with 1-2 years of experience 4, Non-physician
Was a proctor present?	0, No 1, Yes
Was a dual console used?	1, No 2, Yes - Trainee at second console 3, Yes - Consultant at second console 4, Yes - Proctor at second console
Was telepresence (remote input) used?	1, Not available 2, Available, but not used 3, Used - Lead surgeon 4, Used - Second surgeon 5, Used - Proctor 6, Used - Observer
What was the docking time (from first incision to fully docked) in minutes?	Integer
What was the console time (from docking to undocking the robot) in minutes?	Integer
What was the time from skin to skin (incision to closure) in minutes?	Integer
What was the overall theatre time (from patient being wheeled into theatre to being wheeled out of theatre) in minutes?	Integer
What was your port configuration?	1, Robotic-specific layout 2, Same as standard laparoscopic/thoracoscopic or equivalent 3, Single-port 4, Other, please specify
Did you use a wristed stapler?	0, No 1, Yes 99, Unknown
Primary energy source	1, Monopolar and bipolar 2, Advanced bipolar 3, Ultrasonic 4, Combination, please specify 5, Unknown
Was fluorescence imaging (e.g. ICG - indocyanine green) used?	0, No 1, Yes - No change to procedure 2, Yes - Led to a change in the procedure
Did you experience failure of any of the following during the operation?	1, Instrument wrist or tool tip 2, Cautery problem 3, Shaft failure 4, Cable failure 5, Housing failure 6, No failures
Was there additional laparoscopic/thoracoscopic or equivalent assistance required beyond what was planned initially?	0, No 1, Yes
If converted to open (i.e. 20.1 robotic but 21.1 open): What was the reason for conversion?	1, Bleeding 2, Poor exposure 3, Equipment malfunction 4, Progress too slow 5, Oncological concerns 6, Other, please specify
How many other major cases are on the operating theatre list today?	Integer
How many other minor cases are on the operating theatre list today?	Integer
Compared with the approach you would have otherwise used (laparoscopic or equivalent or open), how did the robotic case affect your musculoskeletal comfort?	1, Much worse than alternative 2, Worse than alternative 3, No difference 4, Better than alternative 5, Much better than alternative 99, Unable to answer
Was there a trainee involved in the operative console in this case?	1, No 2, Yes

If abdominal operation (branch from question 11.1 abdomen and pelvis – excluding caesarean section)	
Has the patient had previous abdominal surgery?	0, No 1, Yes, open surgery 2, Yes, minimally invasive surgery 3, Yes, open and minimally invasive surgery
If open in 20.1: Abdominal incision	1, Vertical midline 2, Paramedian 3, Transverse 4, Oblique (e.g. Kocher / McBurney) 5, Flank 6, Pfannenstiel 7, Chevron/Rooftop 8, Thoraco-abdominal 9, Not applicable
If standard minimally invasive or robotic assisted in 20.1: – Was there an extraction site?	0, No 1, Yes - Vertical midline 2, Yes - Paramedian 3, Yes - Transverse 4, Yes - Oblique (e.g. Kocher / McBurney) 5, Yes - Flank 6, Yes - Pfannenstiel 7, Yes - Chevron/Rooftop 8, Yes - Thoraco-abdominal 9, Yes – Other, please specify
Was a stoma created? <i>Please tick all that apply</i>	0, No 1, Yes - end ileostomy 2, Yes – loop ileostomy 3, Yes – double-barrelled ileostomy 4, Yes – end colostomy 5, Yes – loop colostomy 6, Yes – double-barrelled colostomy 7, Yes – urostomy 8, Yes – mucus fistula 9, Other, please specify
Was a bowel anastomosis performed?	1, No 2, Yes – Stapled and intracorporeal 3, Yes – Stapled and extracorporeal 4, Yes – handsewn and intracorporeal 5, Yes – handsewn and extracorporeal
Did any of the following intra-operative events occur? <i>Please tick all that apply</i>	1, Operating time >4h 2, Solid organ injury 3, Ureteric injury 4, Vascular injury 5, Blood transfusion 6, Haemodynamic instability 7, Vasopressor requirement 8, None of the above
Was tranexamic acid used intra-operatively?	0, No 1, Yes - given prophylactically on induction 2, Yes - given in response to bleeding
If there was a colonic resection, did the patient have mechanical bowel preparation pre-operatively	0, No 1, Yes 2, No colonic resection
If clean-contaminated or contaminated or dirty abdominal operation (Q.22.1): Were gloves and instruments changed prior to abdominal wall closure?	0, No 1, Yes 2, Unknown
If open or extraction site: What suture was used for primary fascial closure of the main abdominal incision or extraction site?	1, PDS / PDS II (polydioxanone) 2, PDS Plus (triclosan-coated polydioxanone) 3, Maxon (polyglyconate) 4, Vicryl (polyglactin 910) 5, Vicryl Plus (triclosan-coated polyglactin 910) 6, Vicryl Rapide (irradiated polyglactin 910) 7, Polysorb (lactomer 9-1) 8, Other please specify 9, Unknown – operation note does not record suture material
If open or extraction site: Was the fascial closure technique: <i>Please tick all that apply</i>	1, Continuous running suture 2, Interrupted sutures 3, Small bite technique 4, Large bite technique 5, Unknown
If laparoscopic or robotic assisted abdominal operation: Which port sites were closed at fascial level (not including extraction site)?	0, None 1, All ports sized 5mm and above 2, All ports sized 8mm and above 3, All ports sized 10mm and above
If ports closed: What suture was used for fascial closure of the port sites?	1, PDS / PDS II (polydioxanone) 2, PDS Plus (triclosan-coated polydioxanone) 3, Maxon (polyglyconate) 4, Vicryl (polyglactin 910) 5, Vicryl Plus (triclosan-coated polyglactin 910) 6, Vicryl Rapide (irradiated polyglactin 910) 7, Polysorb (lactomer 9-1) 8, Other please specify 9, Unknown – operation note does not record suture material
If 24.1 skin closed: How was the skin closed? <i>Please tick all that apply</i>	1, Staples 2, Interrupted transcutaneous monocryl 3, Interrupted transcutaneous biosyn 4, Interrupted transcutaneous other, please specify 5, Continuous transcutaneous monocryl 6, Continuous transcutaneous biosyn 7, Continuous transcutaneous other, please specify 8, Interrupted subcuticular monocryl 9, Interrupted subcuticular biosyn 10, Interrupted subcuticular other, please specify 11, Continuous subcuticular monocryl 12, Continuous subcuticular biosyn 13, Continuous subcuticular other, please specify 14, Left open (i.e. skin not closed) or delayed closure 15, Other, please specify 16, Unknown
Did the patient experience any of the following post-operative complications? <i>Please tick all that apply</i>	1, Stoma complications (high output/prolapse/necrosis) 2, Ileus 3, Intra-abdominal collection 4, Fistula (enterocutaneous/enteroenteric) 5, Hernia 6, Acute Kidney injury 7, Electrolyte disturbance 8, Adhesive small bowel obstruction 9, None of the above
If Emergency laparotomy (Branch from Question 9.1 (emergency), 11.1 (abdomen and pelvis), 20.1 (all modalities) and 21.1 (open)) (excluding appendicectomy or cholecystectomy)	
Did the patient have an emergency laparotomy?	0, No 1, Yes
Was the patient seen by or discussed with a Consultant or equivalent surgeon prior to a decision to operate?	0, Not seen or discussed with a Consultant or equivalent surgeon 1, Patient discussed with a Consultant or equivalent surgeon 2, Patient seen by a Consultant or equivalent surgeon
Was the patient seen by or discussed with a Consultant anaesthetist or equivalent prior to the operation?	0, Not seen or discussed with a Consultant or equivalent anaesthetist 1, Patient discussed with a Consultant or equivalent anaesthetist 2, Patient seen by a Consultant or equivalent anaesthetist
Did the patient have a CT scan pre-operatively?	0, No 1, Yes
Was a risk prediction model used before the operation to predict post-operative mortality?	0, No 1, Yes
Did the patient have a history of active cancer in the last 5 years?	0, No 1, Yes
What was the indication for the emergency laparotomy? (Please tick all that apply)	1, Perforation – Oesophagus 2, Perforation – Stomach 3, Perforation – Duodenum 4, Perforation – Small bowel 5, Perforation – Large bowel 6, Perforation – other 7, Sepsis – perforation 8, Sepsis – anastomotic leak 9, Sepsis – abdominal abscess 10, Sepsis – intestinal fistula 11, Sepsis – iatrogenic injury 12, Sepsis – not specified 13, Obstruction - Tender small bowel obstruction 14, Obstruction - Non-tender small bowel obstruction 15, Obstruction - Tender large bowel obstruction 16, Obstruction - Non-tender large bowel obstruction 17, Obstruction - Gastric outlet obstruction 18, Obstruction - Incarcerated/strangulated hernia 19, Obstruction - Hiatus hernia / para-oesophageal hernia 20,

	Obstruction - Volvulus 21, Obstruction - Internal hernia 22, Obstruction - Incisional hernia 23, Obstruction - Intussusception 24, Obstruction - Pseudo-obstruction 25, Obstruction - Foreign body 26, Ischaemia – small bowel 27, Ischaemia – large bowel 28, Ischaemia – stomach 29, Ischaemia – oesophagus 30, Ischaemia – other 31, Other – abdominal wall dehiscence 32, Other – bleeding or haemorrhage 33, Other – abdominal compartment syndrome 34, Other - colitis 35, Other – planned re look laparotomy 36, Other – please specify
Did the patient have a white cell count and/or CRP tested? <i>Please tick all that apply</i>	0, No 1, White cell count 2, CRP
If WCC tested: White cell count ($\times 10^9/L$) (Please record the value closest to the time of decision to operate/booking patient for theatre)	Number
If CRP tested: C-Reactive Protein (CRP) (mg/L) (Please record the value closest to the time of decision to operate/booking patient for theatre)	Number
Did the patient have a blood lactate tested?	1, No - cannot measure in our hospital 2, No - can measure in our hospital, but not performed 3, Yes – arterial blood gas 4, Yes- venous blood gas
If blood lactate: What was the blood lactate (mmol/L)? (If multiple blood lactate readings done please record the value closest to the time of decision to operate/booking patient for theatre)	Number
If adult: Estimated blood loss	1, < 100 ml 2, 100-500 ml 3, 501-1000 ml 4, >1000 ml 5, Not recorded
If child: Estimated blood loss as percentage of estimated blood volume	1, <5% 2, 5-10% 3, 11-20% 4, >20% 5, Not recorded
Was the abdomen left open at the end of the operation (i.e. a damage control laparotomy)?	0, No 1, Yes
If the abdomen was left open: how many more re-look laparotomies did the patient undergo?	Integer
If the abdomen was left open: Which technique was used for the open abdomen at the primary operation?	1, Traditional temporary abdominal closure methods, including but not limited to Bogota bag, Opsite sandwich etc. 2, Vacuum-assisted or negative pressure therapy device 3, Mesh mediated or dynamic fascial closure techniques 4, Skin-only closure options 7, Other, please specify
If the abdomen was left open: What was the closure technique for the open abdomen at the final relook laparotomy?	1, Primary fascial closure 2, Mesh mediated or assisted fascial closure 3, Component separation techniques 4, Bridged fascial closure (when midline cannot be closed) 5, Skin only closure 6, Other, please specify
What was the frequency of post-operative observations for the first 24 hours following the operation? (Observations include blood pressure, heart rate, temperature, oxygen saturations, respiratory rate and conscious level)	0, No observations done in the first 24 hours 1, Observations done between every 0-4 hours 2, More than 4 hours between each set of observations
When were they first reviewed by a Consultant or equivalent surgeon following the operation?	1, Within 6 hours 2, Over 6 hours but less than 12 hours 3, Over 12 hours but less than 24 hours 4, Over 24 hours 5, Not reviewed by a Consultant or equivalent surgeon in the post-operative period
If breast operation (from operative codes Q. 19.1)	
Did the patient have any other following complications? <i>Please tick all that apply</i>	0, No 1, Seroma 2, Haematoma 3, Skin necrosis 4, Nipple necrosis 5, Implant loss 6, Flap necrosis 7, Flap loss

If procedure is Caesarean section (caesarean delivery is an equivalent term) (branch from question 19.1)	
Gravida (number of pregnancies including this pregnancy)	Integer
Parity (number of previous births after 24 weeks gestation)	Integer
Number of previous caesarean sections	Integer
Previous uterine surgery (other than caesarean section)	0, No 1, Yes
Current pregnancy (weeks) <i>For example, if the patient is 38 weeks and 4 days, please enter 38 in this box</i>	Integer
Current pregnancy (days) <i>For example, if the patient is 38 weeks and 4 days, please enter 4 in this box</i>	Integer
Maternal complications during pregnancy <i>Please tick all that apply</i>	1, Anaemia 2, Pre-eclampsia 3, Antepartum haemorrhage 4, Diabetes in pregnancy (gestational) 5, Suspected placenta praevia or placenta accreta spectrum 6, None of the above
Was the woman in labour prior to Caesarean section?	0, No 1, Yes - 1st stage 2, Yes - 2nd stage
Was the labour induced or spontaneous?	1, Induced 2, Spontaneous
Category of urgency	1, Immediate threat to life 2, Maternal or foetal compromise, but not immediately life threatening 3, Needing early delivery but no compromise 4, Elective
If labour occurred (i.e. was the woman in labour prior to Caesarean section? Is 1 or 2): Maternal complications during labour <i>Please tick all that apply</i>	1, Failure induction of labour 2, Placental abruption 3, Delay in first stage 4, Delay in second stage 5, PROM >24 hours 6, Infection (e.g. chorioamnionitis) 7, Failed instrumental birth 8, Obstructed labour 9, Suspected uterine rupture 10, None 11, Other, please specify
Main indication for Caesarean section	1, Failure to progress / labour dystocia 2, Failed induction of labour 3, Suspected foetal compromise 4, Malpresentation 5, Twin pregnancy 6, Previous Caesarean section with contraindication to VBAC 7, Maternal medical conditions 8, Obstructed labour / cephalopelvic disproportion 9, Placenta Praevia (major) & Placenta Accreta spectrum 10, Placenta abruption 11, Suspected uterine rupture 12, Cord prolapse 13, Maternal request 14, Severe foetal macrosomia 15, Other, please specify
Type of skin incision	1, Transverse (Joel Cohen / Pfannenstiel) 2, Midline vertical
Pre-operative haemoglobin (gram/litre)	Integer
Were uterotonics given to prevent post-partum haemorrhage? <i>A uterotonic is a medication that stimulates contraction of the uterine muscles and increases uterine tone, for example oxytocin</i>	0, No 1, Yes
Intra-operative blood loss (millilitres)	Integer
Was blood loss measured or estimated	1, Measured (quantified) 2, Estimated (visual) 3, Mixed (partly measured, partly estimated) 4, Unknown
Did the patient have a post-partum haemorrhage?	0, No 1, Yes - primary post-partum haemorrhage 2, Yes - secondary post-partum haemorrhage 3, Yes - primary and secondary post-partum haemorrhage
Additional blood loss after surgery (ml) up to 24 hours postpartum	Integer
What measures were taken to treat the post-partum haemorrhage? <i>Please tick all that apply</i>	1, Additional uterotonics 2, Artery ligation 3, Balloon tamponade 4, Compression suture (e.g. b-lynnch) 5, Hysterectomy 6, Blood transfusion 7, Tranexamic acid 8, None of the above
Was uterine rupture confirmed at caesarean section?	0, No 1, Yes
Did any of the following intra-operative complications occur? <i>Please tick all that apply</i>	1, Uterine angle extension 2, Impacted foetal head 3, Bladder injury 4, Bowel injury 5, Ureteric injury 6, None of the above
Did the woman have post-operative sepsis, if so, what was the source of this?	0, No 1, Yes - surgical site infection 2, Yes - intra-abdominal sepsis 3, Yes - endometritis 4, Yes - urinary tract 5, Yes - pulmonary source 6, Yes - other, please specify 7, Yes - unknown
Post delivery haemoglobin (gram/litre)	Integer
Did the woman receive a blood transfusion after Caesarean section but before discharge?	0, No 1, Yes
If return to theatre: What was the reason for return to theatre?	1, Repair of wound dehiscence 2, Treatment of post-partum haemorrhage 3, Treatment for sepsis 4, Repair of organ injury 5, Other, please specify
Number of foetuses in pregnancy? <i>(Please complete the below questions for each baby)</i>	1, 1 2, 2 3, 3 4, 4 5, 5 6, 6
Presentation of baby at birth	1, Cephalic 2, Breech 3, Shoulder
Apgar score at 1 minute	Integer
Apgar score at 5 minutes	Integer

Was the baby admitted to the neonatal unit?	0, No 1, Yes
Did the baby suffer any of the following complications within 30 days of birth? <i>Please tick all that apply</i>	1, Birth asphyxia 2, Seizures 3, Skin laceration 4, Cephalohaematoma 5, Skull fracture 6, Brachial plexus injury 7, Neonatal death 8, None 9, Other, please specify

If the patient had a fracture or hand injury (branch from question 10.5 - Single acute fracture / multiple acute fractures / vertebral fracture/hand injury)	
Please identify all the patient's fracture locations. <i>Please tick all that apply</i>	1, Humerus 2, Radius/Ulna 3, Femur 4, Tibia/Fibula 5, Hand 6, Pelvis/Acetabulum 7, Foot 8, Spine 9, Other, please specify
Was this revision surgery?	0, No 1, Yes
Were local antibiotics used during any surgery?	0, No 1, Coated implant 2, Topical powder 3, Local antibiotic carrier e.g. beads, cerament, please specify 4, Irrigation 5, Other, please specify
If humerus fracture: Was this an open fracture of the humerus?	0, No 1, Yes
If radius/ulna fracture: Was this an open fracture of the radius/ulna?	0, No 1, Yes
If femoral fracture: What is the location of the femoral fracture?	31, Proximal 32, Diaphyseal 33, Distal
If 31 Proximal: What kind of fracture is it?	31B (Intracapsular) 31A1/A2 (Extracapsular, Peritrochanteric) 31A3 (Intertrochanteric/Reverse Oblique)
If femoral fracture: Was this an open fracture of the femur?	0, No 1, Yes
If tibia/fibula fracture: What is the location of the tibia/fibula fracture?	41, Proximal 42, Diaphyseal 43, Distal 44, Malleolar segment (Ankle)
If tibia/fibula fracture: Was this an open fracture of the tibia/fibula?	0, No 1, Yes
If hand fracture: What is the location of the hand fracture?	51, Carpal bones 52, Metacarpals 53, Phalanges
If hand fracture: Was this an open fracture of the hand?	0, No 1, Yes
If pelvic/acetabulum fracture: What is the location of the pelvic fracture?	61, Pelvic ring 62, Acetabulum
If pelvic/acetabulum fracture: Was this an open fracture of the pelvis?	0, No 1, Yes
If foot fracture: Was this an open fracture of the foot?	0, No 1, Yes
If spine fracture: Was this an open fracture of the spine?	0, No 1, Yes
If other: Was this an open fracture?	0, No 1, Yes
If it is an ankle fracture (Branch from 44)	
Was there documented peripheral neuropathy?	0, No 1, Yes
Malleolar involvement <i>Please tick all that apply</i>	0, None (isolated syndesmosis injury) 1, Lateral 2, Medial 3, Posterior
Type of definitive fixation <i>Please tick all that apply</i>	1, Plates and/or screws 2, Fibular nail 3, Hindfoot nail 4, External fixator (non-circular) 5, Circular / Hexapod frame 6, Cast 7, Other
Was the syndesmosis fixed? <i>Please tick all that apply</i>	1, No 2, Yes – rigid fixation (screw) 3, Yes – Flexible fixation (tightrope)
If posterior malleolus fracture: Was the posterior malleolus fixed?	0, No 1, Yes
Total time intended restricted or non-weight bearing from surgery? (days)	Integer
Intended immobilisation once unrestricted weight bearing	0, None 1, Cast 2, Boot 3, Other
If hip fracture (Branch from 31B)	
Did the patient receive a hip hemiarthroplasty	0, No 1, Yes
If yes: approach	1. Anterior 2. Anterolateral 3. Posterior 4. Other
If yes: what was the implant?	1. Cemented 2. Uncemented
If cemented: what was the cement?	1. No antibiotic 2. Single antibiotic 3. Dual antibiotic
If hip fracture (Branch from 31A1/2 or 31A3)	
Was the implant	1. Short nail 2. Long nail 3. Sliding hip screw 4. Other
If SHS: was a trochanteric stabilisation plate used?	0, No 1, Yes
If closed tibial fracture (Branch from tib/fib 41, Proximal 42, Diaphyseal 43, Distal and closed)	
Was the fracture fixed with? <i>Please tick all that apply</i>	1, Intramedullary nail 2, Plate and/or screws 3, External fixator (non-circular) 4, Circular / Hexapod frame 5, K wires 6, Amputation
If external fixator was used: was this	1. A temporary fixator 2. Definitive external fixation
If it is an open fracture (branch from any open fracture above) – please complete for each open fracture if polytrauma	

What is the Gustillo-Anderson Classification of the fracture?	1, 1 2, 2 3, 3a 4, 3b 5, 3c
Type of irrigation at debridement	1, Saline only 2, Water 3, Soapy 4, Including betadine 5, Other
What skeletal stabilisation was completed at the time of initial debridement?	1, Definitive stabilisation 2, Provisional stabilisation
Provisional method employed: <i>Please tick all that apply</i>	1, Cast 2, External fixator (non-circular) 3, Circular / Hexapod frame 4, Internal fixation 5, Nothing 6, Removal splint 7, Mono-lateral Exfix + cast
Definitive fixation at initial debridement: <i>Please tick all that apply</i>	1, Amputation 2, Cast 3, External fixator (non-circular) 4, Circular / Hexapod frame 5, Intramedullary nail 6, Plate / screws 7, K wires 8, Mono-lateral ExFix + cast
Definitive method of skeletal stabilisation if not at initial surgery: <i>Please tick all that apply</i>	1, Amputation 2, Cast 3, External fixator (non-circular) 4, Circular / Hexapod frame 5, Intramedullary nail 6, Plate / screws 7, K wires 8, Mono-lateral ExFix + cast
Was a definitive closure achieved at initial debridement?	0, No 1, Yes
If no: what type of wound dressing was used?	1, Negative pressure 2, Standard dressing (not negative pressure) 3, Honey 4, Other, please specify
What was the method employed for definitive closure?	1, Primary closure (at index procedure) 2, Delayed primary closure (i.e. not at index procedure) 3, Skin graft only 4, Local muscle flap (e.g. gastrocnemius, peroneus brevis) 5, Local fasciocutaneous flap (e.g. keystone or propellar) 6, Free flap – fasciocutaneous (ALT, groin flap etc) 7, Free flap – free muscle flap with skin graft (latissimus dorsi, gracilis etc) 8, Left to heal by secondary intention
If primary or delayed closure, was any bone shortening or deformation undertaken	0, No 1, Yes
What was the total number of operations this patient underwent?	Integer
How many days were there between the initial operation and final operation?	Integer
For how long were antibiotics continued following definitive closure (days)	Integer
Hand injury module (Branch from hand injury Q 10.5)	
Please identify the hand injuries present	1, Paediatric nailbed injury (age ≤16 years) 2, Finger laceration with digital nerve injury 3, Other, please specify
If paediatric nailbed injury: "Following repair of the nailbed, was the nail plate re-affixed to the nail bed?"	0, No 1, Yes
If finger laceration with digital nerve injury: Was the digital nerve directly repaired (with sutures)?	0, No 1, Yes
If other: Please specify	Free text

If Ventriculoperitoneal shunt (branch from question 19.1)	
Hydrocephalus aetiology	1, Malformations (e.g., Spina bifida, Chiari) 2, Aqueduct stenosis 3, Post-haemorrhagic (e.g., SAH in adults, neonatal IVH) 4, Tumour – benign 5, Tumour – malignant 6, Trauma 7, Infection 8, Cyst (e.g., colloid, arachnoid, pineal) 9, Idiopathic intracranial hypertension (IIH) 10, Idiopathic Normal Pressure Hydrocephalus (iNPH) 11, Other, please specify
Birth status (up to 18 years old)	1, Pre-term 2, Term 3, Not known
Does this patient have a concurrent CNS infection or within the last 3 months?	0, No 1, Yes
Is this a primary shunt insertion?	0, No 1, Yes
If primary shunt insertion is 'No': Revision shunt insertion (components revised)	1, Ventricular catheter 2, Peritoneal catheter 3, Both 4, Neither (just valve)
What shunt catheters are available in your hospital?	1, Antibiotic only 2, Plain only 3, Both antibiotic and plain
Catheters used for shunt surgery	1, Antibiotic 2, Plain
Shunt infection (30-days)	0, No 1, Yes
Postoperative day shunt infection confirmed (Date of operation is day 0)	Integer
Shunt infection within 12 months: are you happy to be contacted at 12 months to assess for 12-month shunt infection? (By ticking yes, you agree to ensure you have the appropriate regulatory approvals in place and keep the data secure at your hospital (including operation date) and the team running this study module will be in touch closer to the time to assess for 12-month shunt infection)	0, No, please do not contact me 1, Yes, I am happy to be contacted and acknowledge the guidance provided

If Clavien-Dindo V i.e. mortality (branch from question 29.1)	
This form should be completed for patients who die at any time between their operation and postoperative day 30, including during initial admission or after discharge (within 30 days of the operation). The following questions aim to better understand the series of events leading up to this patient's death. Please answer them to the best of your ability, consulting other team members involved in the patient's care where needed. If you are unable to answer the following questions, even with all the information available in your setting, please indicate this as 'Not able to identify a cause of death'. This form should be completed from information and outputs from a morbidity and mortality meeting. If this is not available, then the independent opinion of two senior clinicians should be used. In the event neither of these are possible then the data inputter can complete the form based on the information they have available. Please indicate at the bottom of the form which data source was used.	
On what post-operative day did the patient die? (Day 0 is the day of operation)	Integer
Where did the patient die?	1, In the hospital in which they underwent the index operation 2, In a different hospital to that where they underwent their index operation 3, Out-of-hospital (i.e. at home or in the community) 4, Hospice or equivalent
Was a formal autopsy performed for this patient, and the results available to you?	0, No autopsy performed 1, Autopsy performed, but results not available to me 2, Autopsy performed and results available
Was the case formally reviewed at a departmental Morbidity and Mortality (M&M) meeting or equivalent?	0, No 1, Yes, but the results are not available to me 2, Yes and results available
Cause of death <i>A primary cause is the condition with which the patient presented to health-care services that required surgical treatment (i.e., failure to cure). A cause should be classified as primary if it would have been likely to cause death within 30 days from the disease or presentation requiring surgery.</i> <i>A secondary cause is a complication of the patient's surgical intervention, defined as an unintended or undesired occurrence in the health-care pathway that caused harm to the patient (i.e., failure to rescue)</i>	1, Primary 2, Secondary
Please outline the cause of death If available, please include the cause(s) of death recorded on the death certificate.	Free text
Failure of which organ system using the Advanced Life Support algorithm (ABCDE) predominantly led to this patient's death?	1, Airway, including airway obstruction 2, Breathing, including hypoxia and primary respiratory failure 3, Circulation, including distributive, hypovolaemic, obstructive, cardiogenic and neurogenic shock 4, Disability and exposure, including neurological events, hypo/hyperthermia, toxins, metabolic or electrolyte disturbance 5, Not able to identify a cause of death
Based on the information you have available, was this death preventable?	1, Not preventable 2, Possibly preventable (< 50% likelihood) 3, Probably preventable (>50% likelihood)
Before presentation to hospital, what opportunities or changes in this patient's care may have prevented death? <i>Please tick all that apply</i>	1, Earlier diagnosis or recognition of disease 2, Better access or affordability of primary assessment 3, Better health seeking behaviours 4, More appropriate triage into secondary care or hospital 5, Optimisation of existing long-term conditions 6, Improved physiological reserve at baseline (prehabilitation) 7, Other, please specify 8, None of the above
In the pre-operative phase, but while the patient is in hospital, what opportunities or changes in this patient's care may have prevented death? <i>Please tick all that apply</i>	1, Earlier diagnosis or recognition of disease 2, Better access or affordability of assessment 3, Improved communication between staff members or teams in the hospital 4, Improved monitoring or recognition of deterioration 5, Improved access to diagnostic tests and imaging 6, Earlier initial resuscitative treatment 7, Better availability of resuscitative products (e.g., blood, fluids, drugs) 8, Prompt escalation to senior clinician(s) 9, Earlier decision for surgical intervention 10, More operating theatre capacity or better access to operating theatre 11, Admission under the correct speciality team 12, Incorrect decision for surgery e.g. patient should have been offered a less invasive treatment or non-surgical management or surgery was futile 13, Suboptimal preoperative optimisation; for example lack of recognition of comorbidities, inadequate resuscitation 14, Availability of a higher care environment e.g., high dependency unit or intensive care 15, Improved capacity for non-operative strategies (interventional radiology / endoluminal) 16, Other, please specify 17, None of the above
In the intra-operative phase, what opportunities or changes in this patient's care may have prevented death? <i>Please tick all that apply</i>	1, Improved communication between staff members or teams in the hospital 2, More experienced surgeon present in theatre 3, More experienced anaesthetist present in theatre 4, Improved availability of surgeons from other specialities 5, Better availability of resuscitative products (e.g., blood, fluids, drugs) 6, Avoid surgical technical errors 7, Improved surgical decision making 8, Higher quality equipment or avoid equipment failure 9, More stable supply of electricity 10, Wrong operative decision e.g. colonic anastomosis when should have had stoma 11, Wrong operative approach e.g. open when should be done laparoscopically or reverse 12, Delay in conversion to a different operative approach e.g. laparoscopic converted to open 13, Escalation to senior surgeon 14, Other, please specify 15, None of the above
Following the operation, but whilst the patient is still in hospital, what opportunities or changes in this patient's care may have prevented death? <i>Please tick all that apply</i>	1, Availability of a higher care environment e.g., high dependency unit or intensive care unit 2, Earlier diagnosis or recognition of post-operative complication 3, Improved monitoring or recognition of deterioration 4, Improved access to diagnostic tests and imaging 5, Better availability of resuscitative products (e.g., blood, fluids, drugs) 6, Prompt escalation to senior clinician(s) 7, Earlier decision for surgical, endoscopic or radiological re-intervention 8, More operating theatre capacity or improved access to operating theatre 9, Improved capacity for non-operative rescue strategies (e.g.

	interventional radiology or endoluminal) 10, Improved communication between staff members or teams in the hospital 11, Improved access to rehabilitation 12, Improved symptom relief (pain control, nausea and vomiting) 13, Better access to supplementary nutrition 14, Appropriate venous thromboembolic prophylaxis or anticoagulation 15, Affordability of treatment or care 16, Incorrect decision for reoperation e.g. should have been offered conservative management or less invasive treatment or surgery was futile 17, Other, please specify 18, None of the above					
Following the operation, after the patient had been discharged from hospital, what opportunities or changes in this patient's care may have prevented death? <i>Please tick all that apply</i>	1, Earlier diagnosis or recognition of post-operative complication 2, Improved monitoring or recognition of deterioration 3, Better access or affordability of assessment 4, Better health seeking behaviours 5, More appropriate triage back into secondary care or hospital 6, More regular follow-up with the clinical team 7, Improved access to rehabilitation 8, Improved symptom relief (pain control, nausea and vomiting) 9, Better access to supplementary nutrition 10, Appropriate venous thromboembolic prophylaxis or anticoagulation 11, Affordability of treatment or care 12, Other, please specify 13, None of the above 14, Not discharged from hospital					
Rate to what extent each of the following factors contributed to the patient's death						
Advanced presentation of the disease requiring surgery (e.g. advanced cancer, massive hernia)	1, Not present 2, Present, but did not contribute 3, Contributed slightly 4, Contributed moderately 5, Contributed significantly					
Unstable physiology at the time of presentation (e.g. shock, hypoxia)	1, Not present 2, Present, but did not contribute 3, Contributed slightly 4, Contributed moderately 5, Contributed significantly					
Underlying frailty	1, Not present 2, Present, but did not contribute 3, Contributed slightly 4, Contributed moderately 5, Contributed significantly					
Underlying long-term health conditions, for example diabetes or chronic obstructive pulmonary disease.	1, Not present 2, Present, but did not contribute 3, Contributed slightly 4, Contributed moderately 5, Contributed significantly					
What is the source of the above information? (We would recommend using the outcomes of a morbidity and mortality meeting or two independent senior clinician opinions on the case) <i>Please tick all that apply</i>	0, Data inputter only (including grade) 1, Morbidity and Mortality meeting 2, Two independent senior clinician opinions 3, Other, please specify					
Rate to what extent each of the following possible intraoperative complications contributed to the patient's death <i>Please put a tick on each row of the table</i>		Did not happen	Happened, but did not contribute	Contributed slightly	Contributed moderately	Contributed significantly
Acute respiratory distress syndrome						
Airway injury						
Arrhythmia (including atrial fibrillation)						
Drug reaction / anaphylaxis						
Myocardial infarction						
Organ injury						
Vascular injury						
Pulmonary embolism						
Severe hypotension						
Stroke						
Other, please specify						
Rate to what extent each of the following possible postoperative complications contributed to the patient's death <i>Please put a tick on each row of the table</i>		Did not happen	Happened, but did not contribute	Contributed slightly	Contributed moderately	Contributed significantly
Acute respiratory distress syndrome						
Acute kidney injury						
Anastomotic leak						
Arrhythmia (including atrial fibrillation)						

	Bleeding					
	Deep vein thrombosis					
	Myocardial infarction					
	Pneumonia					
	Pulmonary embolism					
	Stroke					
	Incisional surgical site infection					
	Organ space infection / collection					
	Sepsis other than lower respiratory tract or surgical site infections					
	Other, please specify					
	Rate to what extent each of the following possible underlying chronic conditions contributed to the patient's death <i>Please put a tick on each row of the table</i>		Did not happen	Happened, but did not contribute	Contributed slightly	Contributed moderately
Arrhythmia (including atrial fibrillation)						
Addison's disease						
Asthma						
Chronic heart failure						
Chronic kidney disease						
Chronic liver disease						
Chronic obstructive pulmonary disease						
Cystic fibrosis						
Diabetes						
Dementia						
Epilepsy						
Haematologic al malignancy						
Hypertension						
HIV infection						
Inflammatory bowel disease						
Ischaemic heart disease						
Multiple sclerosis						
Parkinson's disease						
Peripheral arterial disease						

	Solid organ malignancy					
	Stroke or transient ischaemic attack					
	Other chronic lung disease (e.g. pulmonary fibrosis)					
	Other, please specify					

Appendix C: Definitions with referenced question number**4.1 ASA Classification of physical status¹**

Class	Definition	Examples
1	Normal health	Healthy, non-smoking, no or minimal alcohol use
2	Mild systemic disease	Mild diseases only without substantive functional limitations. Examples include (but not limited to): current smoker, social alcohol drinker, pregnancy, 30 < BMI < 40, well-controlled DM/HTN, mild lung disease
3	Severe systemic disease	Substantive functional limitations. One or more moderate to severe diseases. Examples include (but not limited to): poorly controlled DM or HTN, COPD, morbid obesity (BMI ≥ 40), active hepatitis, alcohol dependence or abuse, implanted pacemaker, moderate reduction in ejection fraction, ESRD undergoing regularly scheduled dialysis, premature infant PCA < 60 weeks, history (> 3 months) of MI, CVA, TIA or CAD/stents
4	Severe systemic disease that is a constant threat to life	Examples include (but not limited to): recent (< 3 months) MI, CVA, TIA or CAD/stents, ongoing cardiac ischaemia or severe valve dysfunction, severe reduction in ejection fraction, sepsis, DIC, ARD or ESRD not undergoing regularly scheduled dialysis
5	Moribund: survival not expected without surgery	Examples include (but not limited to): ruptured abdominal/thoracic aneurysm, massive trauma, intracranial bleed with mass effect, ischaemic bowel in the face of significant cardiac pathology or multiple organ/system dysfunction
6	Brain-dead organ donor	

BMI, body mass index; DM, diabetes mellitus; ESRD, end-stage renal disease; HTN, hypertension; PCA, post conceptual age; MI, myocardial infarction; CVA, cerebrovascular accident; TIA, transient

cerebral ischaemic attack; CAD, coronary artery disease; DIC, disseminated intravascular coagulation; ARD, acute respiratory distress.

7.1 Clinical Frailty Scale^{2,3}

Clinical Frailty Scale*



1 Very Fit – People who are robust, active, energetic and motivated. These people commonly exercise regularly. They are among the fittest for their age.



2 Well – People who have **no active disease symptoms** but are less fit than category 1. Often, they exercise or are very **active occasionally**, e.g. seasonally.



3 Managing Well – People whose **medical problems are well controlled**, but are **not regularly active** beyond routine walking.



4 Vulnerable – While **not dependent** on others for daily help, often **symptoms limit activities**. A common complaint is being "slowed up", and/or being tired during the day.



5 Mildly Frail – These people often have **more evident slowing**, and need help in **high order IADLs** (finances, transportation, heavy housework, medications). Typically, mild frailty progressively impairs shopping and walking outside alone, meal preparation and housework.



6 Moderately Frail – People need help with **all outside activities** and with **keeping house**. Inside, they often have problems with stairs and need **help with bathing** and might need minimal assistance (cuing, standby) with dressing.



7 Severely Frail – **Completely dependent for personal care**, from whatever cause (physical or cognitive). Even so, they seem stable and not at high risk of dying (within ~ 6 months).



8 Very Severely Frail – Completely dependent, approaching the end of life. Typically, they could not recover even from a minor illness.



9. Terminally Ill - Approaching the end of life. This category applies to people with **a life expectancy <6 months**, who are **not otherwise evidently frail**.

Scoring frailty in people with dementia

The degree of frailty corresponds to the degree of dementia. Common **symptoms in mild dementia** include forgetting the details of a recent event, though still remembering the event itself, repeating the same question/story and social withdrawal.

In **moderate dementia**, recent memory is very impaired, even though they seemingly can remember their past life events well. They can do personal care with prompting.

In **severe dementia**, they cannot do personal care without help.

* 1. Canadian Study on Health & Aging, Revised 2008.

2. K. Rockwood et al. A global clinical measure of fitness and frailty in elderly people. CMAJ 2005;173:489-495.

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9.2. Sepsis

Sepsis is defined as life-threatening organ dysfunction caused by a dysregulated host response to infection. Organ dysfunction can be identified as an acute change in total SOFA (Sequential Organ Failure Assessment) score ≥ 2 points consequent to the infection.⁴

9.3. Shock

A life-threatening condition of circulatory failure resulting in inadequate tissue perfusion and oxygen delivery, leading to cellular and organ dysfunction. There are 4 types of shock: septic, cardiogenic, hypovolemic and distributive shock. The diagnosis of acute circulatory failure is based on a combination of clinical, hemodynamic and biochemical signs. Clinical findings include hypotension (MAP <65 mmHg), tachycardia, altered mental status, oliguria, and signs of impaired perfusion (e.g., elevated lactate).⁵

10.1. Malignant disease: Also referred to as cancer, malignant tumour or neoplasm; disease characterised by abnormal cells that grow uncontrollably, invade nearby tissues, and may spread to other parts of the body (metastasis).⁴

10.2. Curative surgical procedure: A surgical intervention performed with the intent to completely remove or eradicate a disease (e.g., cancer), aiming for long-term survival or cure.⁵

Palliative surgical procedure: A surgical intervention performed to relieve symptoms, improve quality of life, or manage complications of disease, without the expectation of cure.^{5,6}

14.1. Consultant or equivalent

A senior, fully qualified surgeon with independent responsibility for patient care and clinical decision-making, or a clinician of equivalent seniority and authority within the relevant healthcare system.

(Adapted from: NHS Employers. “The consultant role and responsibilities”; General Medical Council (GMC) guidance on medical roles and responsibilities.)^{9,10}

18.1. Knife to Skin (Start of Operation):

The time point at which the first surgical incision is made in the patient's skin, marking the beginning of the operative procedure.^{11,12}

22.1. Contamination levels^{13,14,15} :

Clean: Respiratory, Genital, GI or GU tracts not entered; no entry into a hollow viscus.

Examples: Hernia repair, elective splenectomy, adhesiolysis, breast surgery, elective hip and knee replacement, abdominal aortic aneurism repair, etc.

Clean-contaminated: Respiratory, Genital, GI or GU tracts entered in controlled condition, controlled entry into a hollow viscus; without spillage of contents of GI/GU tract .

Examples: Appendicectomy (non-inflamed), bowel resection, cholecystectomy (no bile spillage), caesarean section, lung resection, etc.

Contaminated: Minor spillage of contents of GI/GU tracts, incisions in which acute, nonpurulent inflammation is encountered.

Examples: Appendicectomy (inflamed), cholecystectomy (with bile spillage), necrotic bowel resection (no perforation), penetrating trauma, fresh traumatic wound debridement.

Dirty: Gross spillage of contents of GI/GU tracts, peritonitis, established infection.

Examples: Hartmann's procedure for perforated sigmoid colon, laparotomy for perforated gastric ulcer, open fracture, empyema drainage.

28.1. Clavien-Dindo Classification System

Adverse post-operative events may be classified as:

- **Failure of treatment** – This occurs when the original surgery fails to achieve its intended benefits; for example, persistent pain following laparoscopic cholecystectomy or tumour recurrence following cancer surgery.
- **Sequelae:** The recognised consequences of a given procedure; for example, gut malabsorption following a large or small bowel resection or immune deficiency following splenectomy.
- **Complication:** Any deviation from the normal post-operative course that has an adverse effect on the patient and is not either a treatment failure or sequel.

In the Clavien-Dindo classification, the factor determining the severity of a complication is the treatment required¹⁶. Consequently, a given complication may be graded differently depending on how it has been managed. For example, an anastomotic leak may be managed just with antibiotics if it is contained (grade II) or it may require re-operation under anaesthetic (grade III).

Some other considerations:

- Intra-operative complications are not considered unless they have an adverse effect on the patient post-operatively. The only exception to this is intra-operative death; this is classified as grade V.
- All post-operative adverse events are included, even when there is no direct relationship to the surgery.
- All adverse events within the follow-up period (30 days) are included, including following discharge.
- Diagnostic procedures are not included. For example, a diagnostic oesophagoduodenoscopy (OGD) to look for a source of bleeding without any intervention would not be considered a complication, but a therapeutic OGD with clipping of a bleeding vessel would be considered a grade IIIa complication. Since negative exploratory laparotomies are considered to be diagnostic procedures, they should not be recorded as complications.

Clavien -Dindo Grade	Definition (examples listed in <i>italics</i>)
I	Any deviation from the normal postoperative course without the need for pharmacological (other than “allowed therapeutic regimens”), surgical, endoscopic or radiological intervention. Allowed therapeutic regimens are: selected drugs (antiemetics, antipyretics, analgesics, diuretics and electrolyte replacement), physiotherapy and wound infections opened at the bedside but not treated with antibiotics. <i>Examples: Ileus (deviation from the norm); hypokalaemia treated with K; nausea treated with cyclizine; acute kidney injury treated with intravenous fluids.</i>
II	Requiring pharmacological treatment with drugs beyond those allowed for grade I complications; including blood transfusions; total parenteral nutrition. <i>Examples: Surgical site infection treated with antibiotics; myocardial infarction treated medically; deep venous thrombosis treated with enoxaparin; pneumonia or urinary tract infection treated with antibiotics; blood transfusion for anaemia.</i>
III	Requiring surgical, endoscopic or radiological intervention <i>Examples: Therapeutic endoscopic therapy (do not include diagnostic procedures); interventional radiology procedures; return to theatre for any reason</i>
IV	Life-threatening complications requiring critical care management; brain haemorrhage; or ischemic stroke (excluding TIA). <i>Examples: Pneumonia with ventilator support, renal failure with filtration; SAH; stroke</i>
V	Death of a patient

28.2. 30-day Complications:

Surgical site infections

Surgical site infection is defined at 30 days post-surgery using the Centers for Disease Control (CDC) definition¹⁷ of deep incisional or superficial incisional SSI as follows:

1. The infection must occur within 30 days of the index operation
2. The infection must involve the skin, subcutaneous, muscular, or fascial layers of the incision

3. The patient must have at least one of the following: purulent drainage from the wound; organisms detected by wound swab; diagnosed clinically or at imaging; wound opened spontaneously or by a clinician
4. The patient has at least one of the following: pain, tenderness, localized swelling, redness, heat at the wound site, systemic fever ($>38^{\circ}\text{C}$).

Fracture related infection

A fracture-related infection is an infection arising any time after a fracture. The presence of any one of the following features confirms an FRI (confirmatory criteria):

1. Fistula, sinus or wound breakdown
2. Purulent drainage from the wound or presence of pus during surgery
3. Phenotypically indistinguishable pathogens identified by culture from at least two separate deep tissue/implant specimens
4. Presence of microorganisms in deep tissue taken during an operative intervention, as confirmed by histopathological examination.

Suggestive criteria include: skin redness, fever, radiological signs, new onset joint effusion, elevated serum inflammatory markers, persistent, increasing or new onset wound drainage. If suggestive criteria are identified, the diagnosis of FRI should be considered and confirmatory criteria looked for.

Deep organ space infection:

Infection occurring within 30–90 days of surgery involving organs or spaces manipulated during the procedure (deeper than fascia/muscle), defined by purulent drainage from a drain, positive microbiology, or evidence of infection (e.g., abscess) on imaging or examination, meeting criteria for a specific organ/space infection.¹⁷

Postoperative pulmonary complications

Postoperative pulmonary complications will be a secondary outcome. This is a composite of postoperative pneumonia, acute respiratory distress syndrome (ARDS) and unexpected ventilation. This outcome was adapted from the PRISM randomised controlled trial.¹⁸

Postoperative pneumonia

The US Centers for Disease Control (CDC) definition of pneumonia¹⁹ will be used, modified to accommodate limited availability of radiological facilities at some participating centres:

At least **one** of the following:

- Fever ($>38^{\circ}\text{C}$) with no other recognised cause.

- Leucopaenia (white cell count $<4 \times 10^9$) or leucocytosis (white cell count $>12 \times 10^9$).
- For adults >70 years old, altered mental status with no other recognised cause.

AND at least **two** of the following:

- New onset of purulent sputum or change in character of sputum, or increased respiratory secretions, or increased suctioning requirements.
- New onset or worsening cough, or dyspnoea, or tachypnoea.
- Rales, crackles or bronchial breath sounds.
- Worsening gas exchange (hypoxaemia, increased oxygen requirement).

Wherever possible, the diagnosis should be confirmed with a chest radiograph. The following findings confirm pneumonia:

- New or progressive and persistent infiltrates.
- Consolidation.
- Cavitation.

Unexpected ventilation

Unexpected postoperative ventilation was defined as:

- Any episode of non-invasive ventilation, invasive ventilation, or extracorporeal membrane oxygenation after initial extubation after surgery, **or**
- Patient could not be extubated as planned after surgery.

Acute Respiratory Distress Syndrome

Acute Respiratory Distress Syndrome (ARDS) is an acute diffuse, inflammatory lung injury, leading to increased pulmonary vascular permeability, increased lung weight, and loss of aerated lung tissue with hypoxemia and bilateral radiographic opacities. The Berlin consensus definition²⁰ will be used:

Acute Respiratory Distress Syndrome criteria - ALL 4 CRITERIA REQUIRED	
1. Timing	Within 1 week of known clinical insult or worsening respiratory symptoms
2. Chest imaging	Bilateral opacities (not fully explained by effusions / collapse / nodules).
3. Origin	Respiratory failure (not fully explained by cardiac failure / fluid overload).
4. Oxygenation	Mild: $200\text{mmHg} < \text{PaO}_2/\text{FIO}_2 \leq 300\text{mmHg}$ with PEEP or CPAP $\geq 5\text{cm H}_2\text{O}$

	Moderate: $100\text{mmHg} < \text{PaO}_2/\text{FIO}_2 \leq 200\text{mmHg}$ with $\text{PEEP} \geq 5\text{cm H}_2\text{O}$ Severe: $\text{PaO}_2/\text{FIO}_2 \leq 100\text{mmHg}$ with $\text{PEEP} \geq 5\text{cm H}_2\text{O}$ <i>CPAP: continuous positive airway pressure; FIO₂: fraction of inspired oxygen; PaO₂: partial pressure of arterial oxygen; PEEP: positive end-expiratory pressure.</i>
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Pulmonary embolism (PE)

Pulmonary embolism (PE)²¹ is defined as:

- Symptomatic PE confirmed by imaging (computed tomography pulmonary angiogram (CTPA) demonstrating new intraluminal filling defect in a subsegmental or greater sized pulmonary artery; or ventilation/perfusion scanning with a high probability of PE; or pulmonary angiograph demonstrating PE), **or**
- Fatal PE discovered at autopsy or as judged by the clinical team.

Deep vein thrombosis

Deep vein thrombosis (DVT)²² is defined as lower limb deep vein thrombosis with or without symptoms, proven by:

- Lower extremity ultrasonography revealing non-compressibility at the trifurcation of the popliteal vein or above, **or**
- Computed tomography (CT) venography demonstrating a constant intraluminal filling defect above the trifurcation of the popliteal vein.

Pulmonary aspiration or aspiration pneumonitis

Pulmonary aspiration^{23, 24} is the unintentional entry of material from the mouth, pharynx, or gastrointestinal tract into the larynx and lower respiratory tract. Aspirated material may include saliva, food, liquids, oral secretions, vomit, or gastric contents. Depending on the nature and volume of the material aspirated, it may lead to airway obstruction, aspiration pneumonitis, aspiration pneumonia, or acute respiratory distress.

Aspiration pneumonitis^{23, 24} is an acute, non-infectious inflammatory injury to the lungs caused by aspiration of irritating material, most commonly acidic gastric contents. The aspirated material can result in bronchospasm, pulmonary oedema, hypoxaemia and, in severe cases, acute respiratory distress syndrome (ARDS).

Urinary tract infection

A urinary tract infection²⁵ is an event meeting either Symptomatic UTI (SUTI) or Asymptomatic Bacteremic UTI (ABUTI) criteria, specifically:

- a urine culture with no more than two species of organisms, at least one a bacterium at $\geq 100,000$ CFU/mL

combined with either

- (a) at least one qualifying sign or symptom (fever $>38^{\circ}\text{C}$, suprapubic tenderness, costovertebral angle pain/tenderness, urgency, frequency, or dysuria; age-specific criteria for infants ≤ 1 year)
- (b) no such symptoms plus a matching organism in blood.

Central line infection

A central line infection²⁶⁻²⁸ is a laboratory-confirmed bloodstream infection in which a blood stream infection organism is identified and an eligible central line is present on the date of event or date before.

- **Eligible central line:** a central line in place for more than two consecutive calendar days (on or after CL Day 3) following first access in an inpatient location during the current admission; it remains eligible until the day after removal or discharge, whichever comes first.
- **Central line:** an intravascular catheter terminating at, in, or close to the heart or in a great vessel, and used for infusion, blood withdrawal, or haemodynamic monitoring. Includes peripherally placed central lines (e.g. PICC line, Midline)

For sepsis or septic shock (see 9.2 and 9.3).

Postoperative cardiac complications

Acute Myocardial Infarction

Myocardial Infarction (type 1, 2 and 3) as defined by the Fourth Universal Definition of Myocardial Infarction (2018) when 'there is acute myocardial injury with clinical evidence of acute myocardial ischaemia and with detection of a rise and/or fall in cTn values with at least 1 value above the 99th percentile URL and at least one of the following:

- Symptoms of myocardial ischaemia;
- New ischaemic ECG changes;
- Development of pathological Q waves;
- Imaging evidence of new loss of viable myocardium or new regional wall motion abnormalities in a pattern consistent with an ischemic aetiology;

- Identification of a coronary thrombus by angiography or autopsy (not for types 2 or 3 MIs)
- Postmortem demonstration of acute atherothrombosis in the artery supplying the infarcted myocardium meets criteria for *type 1 MI*. Evidence of an imbalance between myocardial oxygen supply and demand unrelated to acute atherothrombosis meets criteria for *type 2 MI*. Cardiac death in patients with symptoms suggestive of myocardial ischaemic and presumed new ischaemic ECG changes before cTN values become available or abnormal meets criteria for *type 3 MI*²⁹.

New acute heart failure³⁰⁻³²

New-onset symptoms and/or signs of heart failure (dyspnoea, orthopnoea, pulmonary crackles, raised JVP, peripheral oedema) arising postoperatively in a patient without a prior diagnosis of heart failure, accompanied by :

- objective evidence of cardiogenic pulmonary or systemic congestion, specifically, at least one of: radiographic pulmonary oedema, echocardiographic evidence of elevated filling pressures or new systolic/diastolic dysfunction, or elevated natriuretic peptide (BNP/NT-proBNP above the guideline threshold for acute presentation)

And

- requiring initiation of treatment (diuretic, vasodilator, inotrope, or non-invasive/invasive ventilatory support).

Clinically significant cardiac arrhythmia requiring treatment³³⁻³⁵

Any new arrhythmia documented on ECG or continuous telemetry that prompts therapeutic intervention: antiarrhythmic or rate-control drug administration, electrical cardioversion or defibrillation, temporary or permanent pacing, or catheter ablation. Arrhythmias managed by observation or by correction of a reversible precipitant alone (e.g. electrolyte repletion) do not meet this criteria.

Stroke³⁶⁻³⁸

An acute episode of focal or global neurological dysfunction caused by brain, spinal cord, or retinal vascular injury, attributable to infarction or haemorrhage. Requires either symptoms/signs persisting ≥24 hours (or until death), or resolution within 24 hours with neuroimaging or pathological confirmation of infarction or haemorrhage in a corresponding vascular territory. Classified as ischaemic, haemorrhagic, or undetermined. Transient ischaemic attack (symptom resolution without imaging evidence of infarction) does not meet this criteria.

Postoperative haemorrhage requiring transfusion³⁹⁻⁴¹

Clinically overt bleeding occurring after the index operation that leads to transfusion of ≥ 1 unit of red cells (or whole blood), and is associated with a fall in haemoglobin not attributable to haemodilution.

Anastomotic leak

The defect of the integrity of a surgical join/anastomosis between two hollow viscera, leading to communication between the intra- and extraluminal compartments.⁴²

Acute kidney injury

Acute Kidney injury⁴³ is defined as any of the following.

- Increase in serum creatinine (SCr) by 0.3mg/dl or more within 48 hours; or
- Increase in SCr to 1.5 times or more of baseline, which is known or presumed to have occurred within the prior 7 days; or
- Urine volume <0.5 ml/kg/h for 6 hours

Wound dehiscence⁴⁴

Partial or complete separation of the layers of a surgical incision after closure, confirmed on clinical examination.

33.1. ERAS (Enhanced Recovery After Surgery): A multimodal, evidence-based perioperative care pathway designed to reduce surgical stress, maintain physiological function, and accelerate recovery after surgery.⁴⁵

Robotics module:

- Robotics Minor and Major Procedures:

Specialty	Minor	Major
General Surgery	Cholecystectomy Inguinal hernia repair	Colectomy Gastrectomy Pancreaticoduodenectomy
Urology	Pyeloplasty Cyst Decortication	Nephrectomy Prostatectomy
Gynaecology	Salpingo-oophorectomy Endometriosis excision	Hysterectomy Sacrocolpopexy

Cardiothoracic	Thymectomy Lung wedge resection	Mitral valve repair CABG Lobectomy
Neurosurgery	Stereotactic brain biopsy SEEG electrode placement	Brain tumour resection Spinal pedicle screw placement
Orthopaedics	Arthroscopy Ligament reconstruction	Total knee arthroplasty Total hip arthroplasty Spinal pedicle screw placement
ENT	Tonsillectomy	Transoral robotic surgery for oropharyngeal tumours Thyroidectomy
Breast	Lumpectomy Sentinel lymph node biopsy	Mastectomy Latissimus dorsi flap breast reconstruction
Plastics	Carpal tunnel release Nerve repair	Lympho-venous anastomosis or another microsurgery procedure Free flap reconstruction
Paediatric	Orchiopexy Pyloromyotomy	Pyeloplasty Fundoplication
Vascular	Varicose vein surgery Venous thrombectomy	Abdominal aortic aneurysm repair Aorto-iliac bypass

The above list is not exhaustive and is intended to provide a small number of examples only.

Abdominal operation module:

- **Gloves and instruments change:** Change of sterile gloves (or outer gloves if double gloved) for the operating surgeons, assistant surgeons, and scrub staff, and instruments including needle holder, forceps and scissors, after completion of the abdominal component of the operation but before handling the wound edges and abdominal closure.⁴⁶
- **Ileus:** Ileus is defined as the “intolerance of oral intake due to inhibition of the gastrointestinal propulsion without signs of mechanical obstruction”. This normally begins three to five days post-operatively and lasts two to three days before resolving.⁴⁷

Emergency Laparotomy module:

- **Laparotomy:** surgical procedure involving an incision through the abdominal wall to gain access to the abdominal cavity. The incision can be made in various locations, including midline, paramedian, transverse and others, depending on the clinical scenario, anatomical site of interest and surgeon's preference. In the case of emergency settings, laparotomy incisions allow rapid and easy access to the peritoneal cavity.⁴⁸

Caesarean delivery module:

- **Post Partum haemorrhage:** Primary postpartum haemorrhage (PPH)⁴⁹ is the most common form of major obstetric haemorrhage. The traditional definition of primary PPH is the loss of 500 ml or more of blood from the genital tract within 24 hours of the birth of a baby after vaginal birth and 1000 ml after caesarean delivery. PPH can be minor (500–1000 ml) or major (more than 1000 ml). Major can be further subdivided into moderate (1001–2000 ml) and severe (more than 2000 ml). Secondary PPH is defined as abnormal or excessive bleeding from the birth canal between 24 hours and 12 weeks postnatally.

- Apgar Score

The Apgar score is a standardised assessment of the clinical status of a newborn immediately after birth, based on five components (colour, heart rate, reflexes, muscle tone, and respiration) each scored from 0 to 2, resulting in a total score ranging from 0 to 10, and reported at 1 and 5 minutes after birth.⁵⁰

Score	0	1	2
Colour	Blue or Pale	Acrocyanotic	Completely Pink
Heart rate	Absent	<100 bpm	>100 bpm
Reflex irritability	No Response	Grimace	Cry or Active Withdrawal
Muscle tone	Limp	Some Flexion	Active Motion
Respiration	Absent	Weak Cry; Hypoventilation	Good, Crying

Fracture module:

- **Gustillo-Anderson Classification**⁵¹

System used to categorise open fractures based on the mechanism of injury, extent of soft tissue damage, and presence of contamination or vascular injury.

The classification divides open fractures into three main types:

- Type I: Clean wound, <1 cm, minimal soft tissue damage.
- Type II: Wound >1 cm, moderate soft tissue injury, no extensive damage, flaps, or avulsions.
- Type III: Extensive soft tissue damage, high-energy trauma, or severe contamination, subdivided into:
 - IIIA: Adequate soft tissue coverage despite extensive injury.
 - IIIB: Extensive soft tissue loss with periosteal stripping and bone exposure, usually requiring flap coverage.
 - IIIC: Associated with arterial injury requiring emergent repair.

References

1. ASA House of Delegates. ASA physical status classification system. 2014.
<https://www.asahq.org/resources/clinical-information/asa-physical-status-classification-system>. (accessed 3/10/2018)
2. Canadian Study on Health & Aging, Revised 2008.
3. K. Rockwood et al. A global clinical measure of fitness and frailty in elderly people. *CMAJ* 2005;173:489-495.
4. Singer M, Deutschman CS, Seymour CW, et al. *The Third International Consensus Definitions for Sepsis and Septic Shock (Sepsis-3)*. *JAMA*. 2016;315(8):801–810.
5. Cecconi M, De Backer D, Antonelli M, et al. *Consensus on circulatory shock and hemodynamic monitoring. Task force of the European Society of Intensive Care Medicine*. *Intensive Care Med*. 2014;40(12):1795–1815.
6. World Health Organization. “Cancer.” WHO Fact Sheets; and DeVita, Hellman, and Rosenberg’s *Cancer: Principles & Practice of Oncology*
7. National Cancer Institute. *NCI Dictionary of Cancer Terms*. Available from:
<https://www.cancer.gov/publications/dictionaries/cancer-terms>
8. World Health Organization. *Palliative care* [Internet]. Available from:
<https://www.who.int/news-room/fact-sheets/detail/palliative-care>
9. NHS Employers. *The consultant role and responsibilities*.

10. General Medical Council. *Good medical practice*. Available from: <https://www.gmc-uk.org/ethical-guidance/ethical-guidance-for-doctors/good-medical-practice>
11. World Health Organization. *WHO Guidelines for Safe Surgery 2009: Safe Surgery Saves Lives*. Geneva: WHO; 2009.
12. Royal College of Surgeons of England. *Good Surgical Practice*. London: RCS Eng; 2014.
13. Yin V, Cobb JP, Wightman SC, Atay SM, Harano T, Kim AW. Centers for Disease Control (CDC) Wound Classification is Prognostic of 30-Day Readmission Following Surgery. *World J Surg*. 2023 Oct;47(10):2392-2400.
14. Prevention of Infection After Gynecologic Procedures: ACOG Practice Bulletin, Number 195. *Obstetrics & Gynecology* 131(6):p e172-e189, June 2018.
15. Chan D et al. Society of Interventional Radiology; Association of periOperative Registered Nurses. Joint Practice Guideline for Sterile Technique during Vascular and Interventional Radiology Procedures: From the Society of Interventional Radiology, Association of periOperative Registered Nurses, and Association for Radiologic and Imaging Nursing, for the Society of Interventional Radiology [corrected] Standards of Practice Committee, and Endorsed by the Cardiovascular Interventional Radiological Society of Europe and the Canadian Interventional Radiology Association. *J Vasc Interv Radiol*. 2012 Dec;23(12):1603-12. doi: 10.1016/j.jvir.2012.07.017. Erratum in: *J Vasc Interv Radiol*. 2013 Apr;24(4):608.
16. Clavien P, Barkun, J, de Oliveira, ML, Vauthey, JN, Dindo, D, Schulick, RD, de Santibañes, E, Pekolj, J, Slankamenac, K, Bassi, C, Graf, R, Vonlanthen, R, Padbury, R, Cameron, JL, Makuuchi, M. The Clavien-Dindo classification of surgical complications: five-year experience. *Annals of surgery*. 2009;250(2):187-96.
17. Centers for Disease Control and Prevention. *National Healthcare Safety Network (NHSN) Patient Safety Component Manual: Surgical Site Infection (SSI) Event*. Atlanta: CDC
18. Pearse, R.M., et al., The Prevention of Respiratory Insufficiency after Surgical Management (PRISM) Trial. Report of the protocol for a pragmatic randomized controlled trial of CPAP to prevent respiratory complications and improve survival following major abdominal surgery. *Minerva Anesthesiol*, 2017. 83(2): p. 175-182.
19. Jain V, Vashisht R, Yilmaz G, et al. Pneumonia Pathology. [Updated 2023 Jul 31]. In: StatPearls [Internet]. Treasure Island (FL): StatPearls Publishing; 2025 Jan-. Available from: <https://www.ncbi.nlm.nih.gov/books/NBK526116/>
20. ARDS Definition Task Force, Ranieri V, Rubenfeld, GD, Thompson, BT, Ferguson, ND, Caldwell, E, Fan, E, Camporota, L, Slutsky, AS. Acute respiratory distress syndrome: the Berlin Definition. *JAMA*. 2012;307(23):2526-33.
21. van der Hulle T, Cheung WY, Kooij S, Beenen LFM, van Bommel T, van Es J, et al. Simplified diagnostic management of suspected pulmonary embolism (the YEARS study): a prospective, multicentre, cohort study. *Lancet*. 2017 Jul 15;390(10091):289-297.
22. Shalhoub J, Lawton R, Hudson J, Baker C, Bradbury A, Dhillon K, et al. Graduated compression stockings as adjuvant to pharmaco-thromboprophylaxis in elective surgical patients (GAPS study): randomised controlled trial. *BMJ*. 2020 May 13;369:m1309.

23. Kollmeier BR, Keenaghan M. Aspiration risk. In: StatPearls [Internet]. Treasure Island (FL): StatPearls Publishing; 2025. Available from: <https://www.ncbi.nlm.nih.gov/books/NBK470169/>
24. MSD Manual Professional Version. Aspiration pneumonitis and pneumonia [Internet]. [cited 2026 Aug 19]. Available from: <https://www.msmanuals.com/professional/pulmonary-disorders/pneumonia/aspiration-pneumonitis-and-pneumonia>
25. Centers for Disease Control and Prevention, National Healthcare Safety Network. Patient Safety Component Manual, Chapter 7: Urinary Tract Infection (Catheter-Associated Urinary Tract Infection [CAUTI] and Non-Catheter-Associated Urinary Tract Infection [UTI]) Events [Internet]. January 2026. Available from: <https://www.cdc.gov/nhsn/pdfs/pscmanual/7psccauticurrent.pdf>
26. Centers for Disease Control and Prevention, National Healthcare Safety Network. Patient Safety Component Manual, Chapter 4: Bloodstream Infection Event (Central Line-Associated Bloodstream Infection and Non-central Line Associated Bloodstream Infection) [Internet]. January 2026. Available from: https://www.cdc.gov/nhsn/pdfs/pscmanual/4psc_clabscurrent.pdf
27. Centers for Disease Control and Prevention, National Healthcare Safety Network. 2026 NHSN Laboratory Confirmed Bloodstream Infection (LCBI) Checklist [Internet]. January 2026. Available from: <https://www.cdc.gov/nhsn/pdfs/checklists/2026-NHSN-Laboratory-Confirmed-Bloodstream-Infection-LCBI-Checklist.pdf>
28. O'Grady NP, Alexander M, Burns LA, Dellinger EP, Garland J, Heard SO, et al. Guidelines for the prevention of intravascular catheter-related infections. Clin Infect Dis. 2011;52(9):1087–99.
29. Thygesen K, Alpert JS, Jaffe AS, Chaitman BR, Bax JJ, Morrow DA, et al. Fourth Universal Definition of Myocardial Infarction (2018). Circulation. 2018;138(20):e618–e51.
30. Bozkurt B, Coats AJS, Tsutsui H, Abdelhamid CM, Adamopoulos S, Albert N, et al. Universal definition and classification of heart failure. Eur J Heart Fail. 2021;23(3):352–80.
31. McDonagh TA, Metra M, Adamo M, Gardner RS, Baumbach A, Böhm M, et al. 2021 ESC Guidelines for the diagnosis and treatment of acute and chronic heart failure. Eur Heart J. 2021;42(36):3599–726.
32. McDonagh TA, Metra M, Adamo M, Gardner RS, Baumbach A, Böhm M, et al. 2023 Focused Update of the 2021 ESC Guidelines for the diagnosis and treatment of acute and chronic heart failure. Eur Heart J. 2023;44(37):3627–39.
33. Beattie WS, Lalu M, Bocock M, Feng S, Wijesundera DN, Nagele P, et al. Systematic review and consensus definitions for the Standardized Endpoints in Perioperative Medicine (StEP) initiative: cardiovascular outcomes. Br J Anaesth. 2021;126(1):56–66.
34. Van Gelder IC, Rienstra M, Bunting KV, Casado-Arroyo R, Caso V, Crijns HJGM, et al. 2024 ESC Guidelines for the management of atrial fibrillation developed in collaboration with EACTS. Eur Heart J. 2024;45(36):3314–414.

35. Zeppenfeld K, Tfelt-Hansen J, de Riva M, Winkel BG, Behr ER, Blom NA, et al. 2022 ESC Guidelines for the management of patients with ventricular arrhythmias and the prevention of sudden cardiac death. *Eur Heart J*. 2022;43(40):3997–4126.
36. Sacco RL, Kasner SE, Broderick JP, Caplan LR, Connors JJ, Culebras A, et al. An updated definition of stroke for the 21st century: a statement for healthcare professionals from the American Heart Association/American Stroke Association. *Stroke*. 2013;44(7):2064–89.
37. Mashour GA, Moore LE, Lele AV, Robicsek SA, Gelb AW. Perioperative care of patients at high risk for stroke during or after non-cardiac, non-neurologic surgery: consensus statement from the Society for Neuroscience in Anesthesiology and Critical Care. *J Neurosurg Anesthesiol*. 2014;26(4):273–85.
38. NeuroVISION Investigators. Perioperative covert stroke in patients undergoing non-cardiac surgery (NeuroVISION): a prospective cohort study. *Lancet*. 2019;394(10203):1022–9.
39. Schulman S, Angerås U, Bergqvist D, Eriksson B, Lassen MR, Fisher W; Subcommittee on Control of Anticoagulation of the Scientific and Standardization Committee of the International Society on Thrombosis and Haemostasis. Definition of major bleeding in clinical investigations of antihemostatic medicinal products in surgical patients. *J Thromb Haemost*. 2010;8(1):202–4.
40. Schulman S, Kearon C; Subcommittee on Control of Anticoagulation of the Scientific and Standardization Committee of the International Society on Thrombosis and Haemostasis. Definition of major bleeding in clinical investigations of antihemostatic medicinal products in non-surgical patients. *J Thromb Haemost*. 2005;3(4):692–4.
41. Mehran R, Rao SV, Bhatt DL, Gibson CM, Caixeta A, Eikelboom J, et al. Standardized bleeding definitions for cardiovascular clinical trials: a consensus report from the Bleeding Academic Research Consortium. *Circulation*. 2011;123(23):2736–47.
42. Rahbari NN, Weitz J, Hohenberge W et al. Definition and grading of anastomotic leakage following anterior resection of the rectum: A proposal by the International Study Group of Rectal Cancer. *Surgery*. 2010; 147(3):339-351
43. KIDGO, KDIGO Clinical Practice Guideline for Acute Kidney Injury, 2012, www.kidney-international.org
44. World Union of Wound Healing Societies. Consensus document: surgical wound dehiscence — improving prevention and outcomes. London: Wounds International; 2018.
45. ERAS® Society. *Guidelines for perioperative care* [Internet]. Available from: <https://erassociety.org>
46. ChEETAh Trial Collaborative. Effect of a sterile glove and instrument change at the time of abdominal wound closure on surgical site infection (ChEETAh): an international, multicentre, cluster randomised trial. *Lancet*. 2022;400:1767–1776.
47. Beach EC, De Jesus O. Ileus. [Updated 2023 Aug 23]. In: StatPearls [Internet]. Treasure Island (FL): StatPearls Publishing; 2025 Jan-. Available from: <https://www.ncbi.nlm.nih.gov/books/NBK558937/>

48. Frassini S, et al. ECLAPTE: Effective Closure of LAParotomy in Emergency-2023 World Society of Emergency Surgery guidelines for the closure of laparotomy in emergency settings. World J Emerg Surg. 2023 Jul 26;18(1):42. doi: 10.1186/s13017-023-00511-w. Erratum in: World J Emerg Surg. 2023 Nov 27;18(1):52. doi: 10.1186/s13017-023-00522-7.
49. WHO. Guideline on prevention and management of PPH.
<https://platform.who.int/docs/default-source/mca-documents/policy-documents/operational-guidance/BRN-MN-32-03-OPERATIONALGUIDANCE-eng-Prevention-Management-PPH.pdf>
50. American Academy of Pediatrics Committee on Fetus and Newborn: American College of Obstetricians and Gynecologists Committee on Obstetric Practice. The Apgar Score. Pediatrics. 2015 Oct;136(4):819-22. doi: 10.1542/peds.2015-2651.
51. Yim GH, Hardwicke JT. The Evolution and Interpretation of the Gustilo and Anderson Classification. J Bone Joint Surg Am. 2018 Dec 19;100(24):e152. doi: 10.2106/JBJS.18.00342. PMID: 30562299.

Appendix D: Statistical Analysis Plan

The primary aim of SurgWeek-2 is to assess complications measured by Clavien-Dindo grade at 30-days following surgery. The secondary outcomes will include 30-day postoperative mortality, 30-day surgical site infection, 30-day postoperative pneumonia, 30-day postoperative pulmonary embolism and deep vein thrombosis, 30-day postoperative cardiac complications, days alive out of hospital at postoperative day 30. Appendix C includes detailed definitions for these outcomes. The analysis will be based on a comparison of complication rates between patients between income status, emergency and elective operating and across specialities.

The study will be conducted according to STROBE (Strengthening the Reporting of Observational Studies in Epidemiology) and SAMPL guidelines (Statistical Analyses and Methods in the Published Literature). Non-parametric data will be summarised with medians and interquartile ranges and differences between groups tested using the Mann-Whitney U test. The χ^2 test will be used for categorical data. Missing data will be included in flowcharts and summary tables, allowing denominators to remain consistent in calculations.

Hierarchical multivariable, mixed-effects logistic regression will be used to explore the associations with the primary and secondary outcome measure, summarised as odds ratios and 95% confidence intervals (C.I.). Clinically plausible patient, disease, operation and location specific factors have been selected a priori for inclusion in adjusted analyses, in order to identify independent predictors of adverse outcomes after surgery.

We anticipate recruitment from approximately 1000 hospitals around the world with a median of 100 patients per hospital (average five mini-teams per hospital and 20 patients collected per mini-team), providing a sample size estimate of 100,000 patients. This is based on previous studies performed by the NIHR Global Health Research Unit on Global Surgery.⁽¹⁻⁴⁾ Analyses will be performed using the R Foundation Statistical Program version 3.1.1 (packages: finalfit, tidyverse).

References:

1. Naluyimbazi R, Fitzgerald TN. Hernia repair as a tracer for elective surgical care. *The Lancet Global Health*. 2024;12(7):e1069-e70.
2. COVIDSurg Collaborative. Elective surgery cancellations due to the COVID-19 pandemic: global predictive modelling to inform surgical recovery plans. *Br J Surg*. 2020;107(11):1440-9.
3. COVIDSurg Collaborative, GlobalSurg Collaborative. Timing of surgery following SARS-CoV-2 infection: an international prospective cohort study. *Anaesthesia*. 2021;76(6):748-58.
4. Glasbey JC, Nepogodiev D, Simoes JFF, Omar O, Li E, Venn ML, et al. Elective Cancer Surgery in COVID-19-Free Surgical Pathways During the SARS-CoV-2 Pandemic: An International, Multicenter, Comparative Cohort Study. *J Clin Oncol*. 2021;39(1):66-78.

Appendix E: Study approvals processes – UK Audit standards

In the UK we would suggest the study is registered as an audit or service evaluation. Within the UK the audit standards we will assess against are the following:

1. World Health Organisation (WHO) Safer Surgery Checklist use
 - a. The WHO Safer Surgery Checklist is one of the only interventions that has been proven to reduce mortality across surgical specialities and in a global context. As such we will assess the use of this in all operations in the UK.⁽¹⁾
 - b. Audit standard: 100%
2. Time to antibiotics in those with sepsis
 - a. Many post-operative deaths can be attributed to sepsis or bleeding.⁽²⁾ In-hospital mortality for those with septic shock approaches 40-60%. Surgical patients presenting with sepsis is common and they will often require source control with surgical interventions.⁽³⁾ However, prior to this the Sepsis 6 bundle advises antibiotics are given within an hour of recognition of sepsis.⁽⁴⁾ National Emergency Laparotomy Audit (NELA) data suggests only 15.4% of patients receive antibiotics within an hour, however we do not have UK wide data for all specialities.⁽⁵⁾
 - b. Therefore, we will assess in those presenting with sepsis who receives intravenous antibiotics within 1 hour of presentation.
 - c. Audit standard: 100%
3. NCEPOD classification of intervention
 - a. The National Confidential Enquiry into Patient Outcome and Death (NCEPOD) was established to in response to concerns about perioperative death. Subsequently they have published numerous reports on this topic. They have created a framework regarding the timing of operative interventions as it is recognised delays in surgery, particularly out of hours is linked with increased morbidity and mortality.⁽⁶⁾
 - b. We will therefore compare emergency admissions time to theatre against the NCEPOD classification entered at the time of booking the patient for theatre.
 - c. Audit standard: 100%

The service evaluation component will describe current clinical practice and assess the uptake and outcomes of robotic surgery in the NHS and describe:

- The availability and use of robotic surgical platforms across participating hospitals
- Characterise patients undergoing robotic surgery
- Compare outcomes (postoperative mortality, complications and reoperation) between surgical approaches (robotic vs open vs other minimally invasive techniques)

This analysis is intended to describe current patterns of practice and benchmark outcomes across surgical approaches, rather than test a new intervention or introduce changes to clinical care.

References:

1. Haynes AB, Weiser TG, Berry WR, Lipsitz SR, Breizat AH, Dellinger EP, et al. A surgical safety checklist to reduce morbidity and mortality in a global population. N Engl J Med. 2009;360(5):491-9.
2. NIHR Global Health Research Unit on Global Surgery. Mechanisms and causes of death after abdominal surgery in low-income and middle-income countries: a secondary analysis of the FALCON trial. Lancet Glob Health. 2024;12(11):e1807-e15.
3. National Institute for Health and Care Excellence. Sepsis 2024 [Available from: <https://cks.nice.org.uk/topics/sepsis/>].
4. Sepsis Trust. Sepsis screening tool acute assessment 2024 [Available from: <https://sepsistrust.org/wp-content/uploads/2024/08/SEPSIS-Trust-Screening-Acute-Tool-Kits-16-2024.pdf>].
5. NELA Audit. Tenth Patient Report of the National Emergency Laparotomy Audit EXECUTIVE SUMMARY 2024 [Available from: <https://rcoa.ac.uk/sites/default/files/documents/2025-09/NELA-ExecSumm2024.pdf>].
6. National confidential enquiry into Patient Outcome and death. The NCEPOD Classification of Intervention 2004 [Available from <https://www.ncepod.org.uk/classification.html>]

NHS Health Research Authority 'is my study research?' tool outcome

University Hospitals Birmingham NHS Foundation Trust Audit approval (CARMS)

Audit title	Surgical outcomes and national/international surgical benchmarks
Audit Code	CARMS-24253

Your audit has now been registered with the Clinical Audit Team and will appear on the [Clinical Audit Registration and Management System](#) (CARMS).

Please ensure that you log on to the system to update it with any progress made. Once your audit is completed you will need to log in to the system and update your audit status to complete. You will also be required to input your 'key' findings and any recommendations (if applicable) that may have come out of the audit.

Please note that all recommendations will be monitored and you be required to update the progress made on each recommendation (action). Also the person nominated for the action will be alerted so ensure you liaise with them appropriately.

This audit code must be quoted to the Medical Records department (ext 8171/8172) or Health Informatics (<http://charon/Requests/>) if you wish to use their services in connection with this audit.

Please only request notes if you will definitely need to view them, as the Medical Records department have limited space and capacity. If you request patient identifiable data from Informatics, this can only be sent to UHB (@uhb.nhs.uk) or nhs.net email accounts.

If you have any queries please contact ClinicalAudit@uhb.nhs.uk.

Thank You

C.A.R.M.S.

Clinical Audit & Registration Management System

Appendix F: REDCap online data collection

Data will be collected and stored online through a secure server running the Research Electronic Data Capture (REDCap) web application. REDCap allows collaborators to enter and store data in a secure system. The REDCap server is managed by the University of Birmingham, UK. Only anonymised data will be uploaded to the database. No patient identifiable data will be collected.

REDCap databases at the University of Birmingham have been successfully used for a number of international studies, including:

- CovidSurg Study (1,040 participating sites across 85 countries), reference: COVIDSurg Collaborative. *Mortality and pulmonary complications in patients undergoing surgery with perioperative SARS-CoV-2 infection: an international cohort study*. Lancet. 2020;396(10243):27-38. doi:10.1016/S0140-6736(20)31182-X.
- European Society of Coloproctology 2017 Left Colon and Rectal Resection Study (335 participating sites across 49 countries), reference: 2017 European Society of Coloproctology (ESCP) Collaborating Group. *The 2017 European Society of Coloproctology (ESCP) international snapshot audit of left colon, sigmoid and rectal resections - Executive Summary*. Colorectal Dis. 2018;20.
- Right Iliac Fossa Treatment Study (290 participating sites across 5 countries), reference: RIFT Study Group on behalf of the West Midlands Research Collaborative. *Identifying children at low-risk of appendicitis: systematic review and prospective, multicentre validation of risk prediction models in children presenting with right iliac fossa pain*. Lancet Child Adolesc Health 2020; 4: 271–80.

The REDCap database used for the SurgWeek-2 study is run by the NIHR Global Health Research Unit on Global Surgery, within the University of Birmingham Virtual Machine architecture which is physically secured. The architecture is the responsibility of the Storage and Virtualisation Team at the University of Birmingham, Edgbaston, Birmingham, B15 2TT. “At rest” encryption is in place on the database server. Raw data will be stored and will remain at the Birmingham site; it will not be offshored to any other location. The site is physically secure. The virtual hosting service is designed to have no single point of failure with physical redundancy deployed for server, network and storage infrastructure. The virtual server software supports live migration of virtual machines between the physical servers called hosts. Live migration is automatically performed to balance the server load across available infrastructure. On physical server failure the virtual machine is automatically restarted on another host. During host maintenance or intrusive maintenance of the virtual server software, virtual machines are manually migrated to prevent any interruption to service. All physical infrastructure is monitored and automatic alerts generated to systems staff on any failure. All virtual machines are not installed on a single physical server but a range of hosts on which virtual machines automatically live migrate on. Therefore, since there is no one physical location for our machine, it can be considered physically secure. All physical infrastructure and the virtual server software are maintained by the University of

Birmingham IT Services. All physical infrastructure is monitored and automatic alerts generated to systems staff on any failure.

The security of the study REDCap database system is governed by the policies of the University of Birmingham UK, in accordance with the requirements of the General Data Protection Regulations (GDPR). The study will be conducted at collaborating sites in accordance with the country-specific data protection requirements. Once data collection is complete, the electronic research files containing anonymised data will be stored on secure non-networked desktop computers for up to 10 years, in line with current regulations. Access will be restricted to the researchers themselves. Personal data will remain securely at local hospitals.

No sensitive or identifiable data will be collected on the database; the patient's clinical team will only upload anonymised data. Access to data will be restricted, each individual collaborator entering data for SurgWeek-2 will have their own username and password. Each patient will be allocated a unique study number at entry. The central research team will not have any access to patient identifiable data. All communication will use this as the identifier. All data will be analysed and reported in summary format. No hospital level data will be reported. No individual will be identifiable. The anonymised data generated by the study will be held centrally at the University of Birmingham UK, and be analysed by Omar Omar (Senior Statistician, University of Birmingham).

Appendix G: Operational Delivery

National leads: This will comprise a network of surgeons and anaesthetists established with previous GlobalSurg and CovidSurg studies. They are responsible for national coordination of the study. Their specific roles will include

- Acting as a link between mini-teams / Hospital Leads, and the steering committee
- Active engagement with dissemination of the study and other NIHR Unit on Global Surgery Collaborative activities in their country.
- Effective and responsive communication with the steering committee, and with local collaborators
- Recruitment of hospitals within their country
- Registering hospital leads and their teams
- Securing national ethical or regulatory approvals if required

Hospital Lead: This will be a single lead point of contact for data collection at each site who has overall responsibility for site governance registration and coordinating handover between local collaborator teams. They should be a consultant or senior clinician. Minimum requirements for authorship on this study as a Hospital Lead include

- Obtaining local approvals for conduct of the study (e.g. registration of the audit, seeking Caldicott guardian permission to upload data to REDCap, submitting the protocol to Ethics Commission where applicable)
- Coordination of handover between all local collaborator teams at the centre
- Identifying Hospital co-leads
- Involvement in local dissemination and presentation of local results at their centre from the study (or otherwise arranges another collaborator to present on their behalf)

Hospital co-leads: This will be a lead point of contact for data collection in each hospital. They will support the Hospital lead to identify local collaborators to be in the mini-teams and support the mini-teams to identify patients and collect data. The minimum requirements for authorship on outputs of this cohort study include:

- Compliance with local audit approval processes and data governance policies.
- Active involvement in mini-team recruitment and engagement
- Collaboration with the hospital lead to ensure that the results are reported back to the audit office / clinical

Local Collaborators (Data Collectors): This will comprise a team of up to 3 people responsible for data collection over a week period within a single or multiple specialities within their hospital. The ideal team would be a heterogeneous group at different levels of clinical training (e.g. medical students, trainees, consultants). The minimum requirements for authorship on outputs of this cohort study include:

- Compliance with local audit approval processes and data governance policies.
- Active involvement in data collection and uploading it to REDCap
- Collaboration with the regional / local lead to ensure that the results are reported back to the audit office / clinical